# => D HIS

L3

(FILE 'HOME' ENTERED AT 18:29:34 ON 16 MAR 2006)

FILE 'REGISTRY' ENTERED AT 18:29:49 ON 16 MAR 2006

L10 S DOCUSATE/CN L2

3 S DOCUSATE

FILE 'CAPLUS' ENTERED AT 18:32:43 ON 16 MAR 2006

41 S DOCUSATE AND QUATERNARY AMMONIUM

L423 S L3 AND PY<2002

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=> s docusate/cn
             0 DOCUSATE/CN
=> s docusate
             3 DOCUSATE
=> d
L2
     ANSWER 1 OF 3 REGISTRY COPYRIGHT 2006 ACS on STN
RN
     7491-09-0 REGISTRY
ED
     Entered STN: 16 Nov 1984
    Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, potassium salt
           (CA INDEX NAME)
OTHER CA INDEX NAMES:
     Succinic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, potassium salt (8CI)
OTHER NAMES:
CN
    Bis(2-ethylhexyl) potassium sulfosuccinate
CN
    Bis-2-ethylhexyl-2-sulfobutane-1,4-dioate potassium salt
    Dioctyl potassium sulfosuccinate
CN
CN
    Docusate potassium
    Potassium bis(2-ethylhexyl) sulfosuccinate
CN
CN
    Potassium dioctyl sulfosuccinate
CN
    Rectalad Enema
DR
    170717-32-5
    C20 H38 O7 S . K
MF
     STN Files: BEILSTEIN*, BIOTECHNO, CA, CAPLUS, CHEMLIST, CIN, CSCHEM,
LC
      DIOGENES, EMBASE, IFICDB, IFIPAT, IFIUDB, MRCK*, PS, TOXCENTER, USAN,
      USPAT2, USPATFULL
         (*File contains numerically searchable property data)
     Other Sources:
                     EINECS**
         (**Enter CHEMLIST File for up-to-date regulatory information)
CRN
     (10041-19-7)
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K

=> d 12 2-3 ibib abs hitstr

54 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

54 REFERENCES IN FILE CAPLUS (1907 TO DATE)

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The following are valid formats:

Substance information can be displayed by requesting individual fields or predefined formats. The predefined substance formats are: (RN = CAS Registry Number)

REG - RN

SAM - Index Name, MF, and structure - no RN
```

SAM - Index Name, MF, and structure - no RN
FIDE - All substance data, except sequence data
IDE - FIDE, but only 50 names
SQIDE - IDE, plus sequence data

'IBIB' IS NOT A VALID FORMAT FOR FILE 'REGISTRY'
'ABS' IS NOT A VALID FORMAT FOR FILE 'REGISTRY'
'HITSTR' IS NOT A VALID FORMAT FOR FILE 'REGISTRY'

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ANSWER 2 OF 3 REGISTRY COPYRIGHT 2006 ACS on STN
L2
RN
     577-11-7 REGISTRY
ED
     Entered STN: 16 Nov 1984
CN
     Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, sodium salt (9CI)
     (CA INDEX NAME)
OTHER CA INDEX NAMES:
     Aerosol OT-B (6CI)
OTHER NAMES:
CN
     1,4-Bis(2-ethylhexyl) sodium sulfosuccinate
CN
     Adekacol EC 8600
CN
     Aerosol A 501
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     Aerosol AOT
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     Aerosol OT 94
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     Aerosol OT-A
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     Aerosol OT-S
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CN
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     Complemix
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     Constonate
CN
     Coprol
CN
     Coprola
CN
     Correctol Stool Softener Laxative
CN
     Defilin
CN
     DESS
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     Di(2-ethylhexyl) sulfosuccinate sodium salt
CN
     Di-2-ethylhexyl sodium sulfosuccinate
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     Dioctlyn
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CN
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ADDITIONAL NAMES NOT AVAILABLE IN THIS FORMAT - Use FCN, FIDE, or ALL for
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CI
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LC
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BIOTECHNO, CA, CABA, CAOLD, CAPLUS, CASREACT, CBNB, CHEMCATS, CHEMINFORMRX, CHEMLIST, CIN, CSCHEM, CSNB, DDFU, DETHERM\*, DIOGENES, DRUGU, EMBASE, GMELIN\*, HSDB\*, IFICDB, IFIPAT, IFIUDB, IMSCOSEARCH, IPA, MRCK\*, MSDS-OHS, NIOSHTIC, PIRA, PROMT, PROUSDDR, PS, RTECS\*, SCISEARCH, TOXCENTER, USAN, USPAT2, USPATFULL, VETU

(\*File contains numerically searchable property data)
Other Sources: DSL\*\*, EINECS\*\*, TSCA\*\*, WHO

(\*\*Enter CHEMLIST File for up-to-date regulatory information)

CRN (10041-19-7)

Na

#### \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

8049 REFERENCES IN FILE CA (1907 TO DATE)

47 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

8067 REFERENCES IN FILE CAPLUS (1907 TO DATE)

16 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

L2 ANSWER 3 OF 3 REGISTRY COPYRIGHT 2006 ACS on STN

RN 128-49-4 REGISTRY

ED Entered STN: 16 Nov 1984

CN Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, calcium salt (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN Succinic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, calcium salt (8CI) OTHER NAMES:

CN Bis(2-ethylhexyl) calcium sulfosuccinate

CN Bis(2-ethylhexyl) sulfosuccinic acid calcium salt

Calcium bis(2-ethylhexyl) sulfosuccinate

CN Calcium di-2-ethylhexyl sulfosuccinate

CN Calcium dioctyl sulfosuccinate

CN Dioctyl calcium sulfosuccinate

CN Docusate calcium

CN Doxical

CN

CN

MF

CRN

Sulfosuccinic acid, bis(2-ethylhexyl) ester, calcium salt

CN Surfak

C20 H38 O7 S . 1/2 Ca

LC STN Files: ADISNEWS, BEILSTEIN\*, BIOSIS, BIOTECHNO, CA, CAPLUS, CHEMLIST, CIN, DDFU, DIOGENES, DRUGU, EMBASE, IFICDB, IFIPAT, IFIUDB, IPA, MRCK\*, PROMT, PS, TOXCENTER, USAN, USPAT2, USPATFULL (\*File contains numerically searchable property data)

Other Sources: DSL\*\*, EINECS\*\*

(\*\*Enter CHEMLIST File for up-to-date regulatory information)
(10041-19-7)

110 REFERENCES IN FILE CA (1907 TO DATE)
110 REFERENCES IN FILE CAPLUS (1907 TO DATE)

ANSWER 7 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN T.4 ACCESSION NUMBER: 2000:608551 CAPLUS DOCUMENT NUMBER: 133:213151 TITLE: Pharmaceutical compositions and methods for improved delivery of hydrophobic therapeutic agents INVENTOR(S): Patel, Manesh V.; Chen, Feng-Jing PATENT ASSIGNEE(S): Lipocine, Inc., USA SOURCE: PCT Int. Appl., 98 pp. CODEN: PIXXD2 DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: 13 PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO.

WO 20000500	07	A1	20000831	WO 2000-US165	20000105 /
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				GB, GD, GE, GH, GM,	
IN,	IS, JP,	KE,	KG, KP, KR,	KZ, LC, LK, LR, LS,	LT, LU, LV, MA,
MD,	MG, MK,	MN,	MW, MX, NO,	NZ, PL, PT, RO, RU,	SD, SE, SG, SI,
				UA, UG, UZ, VN, YU,	
				SZ, TZ, UG, ZW, AT,	
				IT, LU, MC, NL, PT,	
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			20040401		20000103 <
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JP 20025373	17	T2	20021105	JP 2000-600619	20000105
NZ 513810		Α	20040227	NZ 2000-513810	20000105
RITY APPLN.				US 1999-258654	
				WO 2000-US165	
The procent	4		1.5	iglygoride from phor	·

DATE

The present invention relates to triglyceride-free pharmaceutical compns. AB for delivery of hydrophobic therapeutic agents. Compns. of the present invention include a hydrophobic therapeutic agent and a carrier, where the carrier is formed from a combination of a hydrophilic surfactant and a hydrophobic surfactant. Upon dilution with an aqueous solvent, the composition forms a clear, aqueous dispersion of the surfactants containing the therapeutic agent. The invention also provides methods of treatment with hydrophobic therapeutic agents using these compns. A pharmaceutical composition contained cyclosporin 0.14, Cremophor RH-40 0.41, Arlacel186 0.29, sodium taurocholate 0.26, and propylene glycol 0.46 mg.

REFERENCE COUNT: THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT ANSWER 6 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2001:31306 CAPLUS

DOCUMENT NUMBER: 134:105846

TITLE: Clear aqueous dispersions of triglycerides and

surfactants for delivery of drugs and nutrients

INVENTOR(S): Chen, Feng-Jing; Patel, Mahesh V.

PATENT ASSIGNEE(S): Lipocine, Inc., USA SOURCE: PCT Int. Appl., 103

PCT Int. Appl., 103 pp. CODEN: PIXXD2

CODEN: PIA

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 13

PATENT INFORMATION:

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KIND DATE APPLICATION NO.
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                       A1 20010111 WO 2000-US15133
    WO 2001001960
                                                            20000602 <--
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           CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU,
            ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU,
           LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD,
            SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW
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PRIORITY APPLN. INFO.:
                                         US 1999-345615
                                                           A 19990630
                                         WO 2000-US15133
                                                           W 20000602
```

The present invention relates to drug and nutrient delivery systems, and AB in particular to pharmaceutical compns. and methods for improved solubilization of triglycerides and improved delivery of therapeutic agents. Compns. of the present invention include a triglyceride and a carrier, where the carrier is formed from a combination of at least two surfactants, at least one of which is hydrophilic. Upon dilution with an aqueous solvent, the composition forms a clear, aqueous dispersion of the triglyceride and surfactants. An optional therapeutic agent can be incorporated into the composition, or can be co-administered with the composition. The invention also provides methods of enhancing triglyceride solubility and methods of treatment with therapeutic agents using these compns. Several formulations were presented of compns. that can be prepared according to the present invention using a variety of therapeutic agents. Examples of aqueous dispersions include: (1) Cremophor RH-40 0.75, Peccol 0.25, corn oil 0.40, and fenofibrate 0.10; (2) Cremophor RH-40 0.57, Crovol M-40 0.43, corn oil 0.40, and Rofecoxib 0.15; (3) Tween 80 0.70, Tween 85 0.35, Miglyol 812 0.30, Paclitaxel 0.10, and PEG 400 0.25; or (4) Kessco PEG 400 MO 0.33, corn oil 0.30, and Terbinafine 0.25 parts, resp.

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 12 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2000:123188 CAPLUS DOCUMENT NUMBER: 132:171126 TITLE: Flocculated suspension of megestrol acetate INVENTOR(S): Ragunathan, Narayan; Chao, James C.; Femia, Robert A.; Ross, Malcolm S. F. PATENT ASSIGNEE(S): Pharmaceutical Resources, Inc., USA SOURCE: U.S., 5 pp. CODEN: USXXAM DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. ---------------------20000222 US 1998-63241 US 6028065 Α 19980420 <--WO 2001026626 20010419 WO 1999-US23340 A1 19991007 <--W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG AU 9962942 20010423 AU 1999-62942 **A1** 19991007 <--US 1999-416841 US 6268356 В1 20010731 19991012 <--US 2001048931 US 2001-757261 20010109 <--A1 20011206 US 6593318 B2 20030715 US 2002173497 **A1** 20021121 US 2002-136823 20020430 US 6593320 B2 20030715 PRIORITY APPLN. INFO.: US 1998-63241 A 19980420 A 19991007 WO 1999-US23340 US 1999-416841 A1 19991012 US 2001-757261 A3 20010109 A novel oral antineoplastic composition comprises a stable flocculated AB suspension in water containing megestrol acetate, ≥1 of PEG, propylene glycol, glycerol, and sorbitol, and a surfactant, provided polysorbate and PEG are not simultaneously present. Any surfactant, regardless of the

AB A novel oral antineoplastic composition comprises a stable flocculated suspension in water containing megestrol acetate, ≥1 of PEG, propylene glycol, glycerol, and sorbitol, and a surfactant, provided polysorbate and PEG are not simultaneously present. Any surfactant, regardless of the length of the hydrophobic contact area on its hydrophobic group, can effectively wet megestrol acetate and form a stable flocculated suspension, provided ≥1 of the other named compds. is present. Thus, a suspension was prepared containing megestrol acetate 4.000, glycerol 5.000, sorbitol 15.000, docusate Na (surfactant) 0.002, xanthan gum 0.250, NaOBz 0.200, citric acid 0.300, Na citrate 0.060, sucrose 5.000, lemon flavoring 0.080, and H2O 70.108 weight%.

REFERENCE COUNT: 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> d

L1 HAS NO ANSWERS

L1

STR

Structure attributes must be viewed using STN Express query preparation.

=> s l1 full

#### REG1stRY INITIATED

Substance data SEARCH and crossover from CAS REGISTRY in progress... Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

FULL SEARCH INITIATED 14:59:34 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 5534 TO ITERATE

100.0% PROCESSED 5534 ITERATIONS

3981 ANSWERS

SEARCH TIME: 00.00.01

L2 3981 SEA SSS FUL L1

L3 13736 L2

=> s 13 and ( quaternary ammonium or quaternary phosphonium)

124926 QUATERNARY

361380 AMMONIUM

61727 QUATERNARY AMMONIUM

(QUATERNARY (W) AMMONIUM)

124926 QUATERNARY

15672 PHOSPHONIUM

1179 QUATERNARY PHOSPHONIUM

(QUATERNARY (W) PHOSPHONIUM)

L4 701 L3 AND ( QUATERNARY AMMONIUM OR QUATERNARY PHOSPHONIUM)

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188 DOCUSATE

L5 41 L4 AND DOCUSATE

=> s 15 and py<2002

L6

21808282 PY<2002

23 L5 AND PY<2002

=> s 16 and sulfosuccinic acid

2002 SULFOSUCCINIC

4114809 ACID

1906 SULFOSUCCINIC ACID

(SULFOSUCCINIC (W) ACID)

L7 2 L6 AND SULFOSUCCINIC ACID

### => d 1-2 ibib abs hitstr

ANSWER 1 OF 2 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:652447 CAPLUS

DOCUMENT NUMBER: 141:179653

TITLE: Novel nimesulide compositions

INVENTOR(S): Bosch, H. William; Wertz, Christian F. PATENT ASSIGNEE(S): Elan Pharma International Ltd., USA

SOURCE: U.S. Pat. Appl. Publ., 27 pp., Cont.-in-part of U.S.

Ser. No. 276,400. CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 16

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2004156872	A1	20040812	US 2003-697703	20031031
US 6316029	B1	20011113	US 2000-572961	20000518 <
US 2004013613	A1	20040122	US 2003-276400	20030115
PRIORITY APPLN. INFO.:			US 2000-572961	A1 20000518
			US 2003-276400	A2 20030115
			WO 2001-US15983	W 20010518

The present invention provides nanoparticulate nimesulide compns. The compns. preferably comprise nimesulide and at least one surface stabilizer adsorbed on or associated with the surface of the nimesulide particles. The nanoparticulate nimesulide particles preferably have an effective average particle size of less than about 2000 nm. The composition further comprises one or more addnl. compds., e.g., an analgesic, an anti-inflammatory agent, an antipyretic, a vasomodulator, etc. The invention also provides methods of making and using nanoparticulate nimesulide compns. For example, nimesulide nanoparticles were prepared by combining 0.85 g of Plasdone S-630 dissolved in 79.9 g of water (1% weight/weight) as a surface stabilizer with 4.25 g nimesulide (5% weight/weight) and PolyMill-200 Polystyrene Milling Media and milling for 1 h at 4200 rpm with chilled water (10°) recirculated through the milling chamber. The process yielded a colloidal dispersion of nimesulide with a mean particle size of 150 nm, a D50 of 124 nm, a D90 of 256 nm, and a D95 of 293 nm.

577-11-7, Docusate sodium 5138-18-1D,

Sulfosuccinic acid, dialkyl esters, sodium salts

10041-19-7, Dioctyl sulfosuccinate

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (nimesulide nanoparticulate compns. comprising surface stabilizer)

RN 577-11-7 CAPLUS

CN Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, sodium salt (9CI) (CA INDEX NAME)

Na

RN 5138-18-1 CAPLUS

CN Butanedioic acid, sulfo- (9CI) (CA INDEX NAME)

$$^{\mathrm{SO_3H}}_{|}$$
  $^{\mathrm{HO_2C-CH-CH_2-CO_2H}}_{|}$ 

RN 10041-19-7 CAPLUS CN 'Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester (9CI) (CA INDEX NAME)

7 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2000:123188 CAPLUS

DOCUMENT NUMBER: 132:171126

TITLE: Flocculated suspension of megestrol acetate

INVENTOR(S): Ragunathan, Narayan; Chao, James C.; Femia, Robert A.;

Ross, Malcolm S. F.

PATENT ASSIGNEE(S): Pharmaceutical Resources, Inc., USA

SOURCE: U.S., 5 pp. CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.			
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KG, KP, KR,	KZ, LC, LK, LR,	LS, LT, LU, LV, MD, MG	, MK, MN, MW,		
MX, NO, NZ,	PL, PT, RO, RU,	SD, SE, SG, SI, SK, SL	, TJ, TM, TR,		
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CG, CI, CM,	GA, GN, GW, ML,	MR, NE, SN, TD, TG	,,,		
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		US 2002-136823	20020420		
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PRIORITY APPLN. INFO.:	B2 20030713		3 10000100		
PRIORITI APPEN. INFO.:		US 1998-63241			
			A 19991007		
			A1 19991012		
AB A novel oral antine		US 2001-757261	A3 20010109		

AB A novel oral antineoplastic composition comprises a stable flocculated suspension in water containing megestrol acetate, ≥1 of PEG, propylene glycol, glycerol, and sorbitol, and a surfactant, provided polysorbate and PEG are not simultaneously present. Any surfactant, regardless of the length of the hydrophobic contact area on its hydrophobic group, can effectively wet megestrol acetate and form a stable flocculated suspension, provided ≥1 of the other named compds. is present. Thus, a suspension was prepared containing megestrol acetate 4.000, glycerol 5.000, sorbitol 15.000, docusate Na (surfactant) 0.002, xanthan gum 0.250, NaOBz 0.200, citric acid 0.300, Na citrate 0.060, sucrose 5.000, lemon flavoring 0.080, and H2O 70.108 weight%.

IT 577-11-7, Docusate sodium

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (surfactant; flocculated suspension of megestrol acetate)

RN 577-11-7 CAPLUS

CN Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, sodium salt (9CI) (CA INDEX NAME)

Na

IT 5138-18-1D, Sulfosuccinic acid, esters with

fatty alcs.

CN

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (surfactants; flocculated suspension of megestrol acetate)

RN 5138-18-1 CAPLUS

Butanedioic acid, sulfo- (9CI) (CA INDEX NAME)

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REFERENCE COUNT: 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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_	u	***	_

	(FILE 'HOME' ENTERED AT 14:58:23 ON 16 MAR 2006)
L1	FILE 'CAPLUS' ENTERED AT 14:58:47 ON 16 MAR 2006 STRUCTURE UPLOADED S L1
L2	FILE 'REGISTRY' ENTERED AT 14:59:34 ON 16 MAR 2006 3981 S L1 FULL
	ETTE 'CADITIS' ENTEDED AT 14.50.35 ON 16 MAD 2006

FILE 'CAPLUS' ENTERED AT 14:59:35 ON 16 MAR 2006 13736 S L2 FULL

L3 L4

701 S L3 AND ( QUATERNARY AMMONIUM OR QUATERNARY PHOSPHONIUM) 41 S L4 AND DOCUSATE

L5L6

23 S L5 AND PY<2002

2 S L6 AND SULFOSUCCINIC ACID

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L7

L6 ANSWER 1 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:652447 CAPLUS

DOCUMENT NUMBER: 141:179653

TITLE: Novel nimesulide compositions

INVENTOR(S): Bosch, H. William; Wertz, Christian F. PATENT ASSIGNEE(S): Elan Pharma International Ltd., USA

SOURCE: U.S. Pat. Appl. Publ., 27 pp., Cont.-in-part of U.S.

Ser. No. 276,400. CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	ΑP	PLICATION NO.		DATE	
US 2004156872	<b>A</b> 1	20040812	US	2003-697703		20031031	
us 6316029	B1	20011113	US	2000-572961		20000518	<
US 2004013613	A1	20040122	US	2003-276400		20030115	
PRIORITY APPLN. INFO.:			· US	2000-572961	A1	20000518	
			US	2003-276400	A2	20030115	
			WO	2001-US15983	W	20010518	

AB The present invention provides nanoparticulate nimesulide compns. The compns. preferably comprise nimesulide and at least one surface stabilizer adsorbed on or associated with the surface of the nimesulide particles. The nanoparticulate nimesulide particles preferably have an effective average particle size of less than about 2000 nm. The composition further comprises one or more addnl. compds., e.g., an analgesic, an anti-inflammatory agent, an antipyretic, a vasomodulator, etc. The invention also provides methods of making and using nanoparticulate nimesulide compns. For example, nimesulide nanoparticles were prepared by combining 0.85 g of Plasdone S-630 dissolved in 79.9 g of water (1% weight/weight) as a surface stabilizer with 4.25 g nimesulide (5% weight/weight) and PolyMill-200 Polystyrene Milling Media and milling for 1 h at 4200 rpm with chilled water (10°) recirculated through the milling chamber. The process yielded a colloidal dispersion of nimesulide with a mean particle size of 150 nm, a D50 of 124 nm, a D90 of 256 nm, and a D95 of 293 nm.

IT 577-11-7, Docusate sodium 5138-18-1D,

Sulfosuccinic acid, dialkyl esters, sodium salts 10041-19-7,

Dioctyl sulfosuccinate

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (nimesulide nanoparticulate compns. comprising surface stabilizer)

RN 577-11-7 CAPLUS

CN Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, sodium salt (9CI) (CA INDEX NAME)

Na

RN 5138-18-1 CAPLUS

CN Butanedioic acid, sulfo- (9CI) (CA INDEX NAME)

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RN 10041-19-7 CAPLUS

CN Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester (9CI) (CA INDEX NAME)

L6 ANSWER 2 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2001:780648 CAPLUS

DOCUMENT NUMBER:

135:335147

TITLE:

Polymer-based injectable sustained release

pharmaceutical compositions for peptide and protein

drugs

INVENTOR(S):

Lee, Hee-yong; Lee, Hye-suk; Kim, Jung-soo; Kim,

Sang-beom; Lee, Ji-suk; Choi, Ho-il; Chang, Seung-gu

PATENT ASSIGNEE(S):

Peptron Inc., S. Korea

SOURCE:

PCT Int. Appl., 37 pp.

DOCUMENT TYPE:

CODEN: PIXXD2

DOCUMENT I

Patent English

LANGUAGE:

1 Flights

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION N

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								KR 2	000-	4934	4		A 2	0000	824
							1	WO 2	001-	KR462	2	Ī	w 2	0010	322
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INFO.:	1078687 A1 20011025 WO 20 AE, AG, AL, AM, AT, AU, AZ, BA, BB, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, HU, ID, IL, IN, IS, JP, KE, KG, KP, LV, MA, MD, MG, MK, MN, MW, MX, MZ, SE, SG, SI, SK, SL, TJ, TM, TR, TT, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, DE, DK, ES, FI, FR, GB, GR, IE, IT, BJ, CF, CG, CI, CM, GA, GN, GW, ML, 1099583 A 20011109 KR 20 A1 20020320 EP 20 AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, SI, LT, LV, FI, RO 3026844 A1 20030206 US 20 PLN. 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INFO.:  KR 2000—	1078687 A1 20011025 W0 2001-KR463 AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, HU, ID, IL, IN, IS, JP, KE, KG, KP, KZ, LC, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM : GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, 1099583 A 20011109 KR 2000-49344 A1 20020320 EP 2001-91789 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, IE, SI, LT, LV, FI, RO 3026844 A1 20030206 US 2002-18870 PLN. INFO.:  KR 2000-20484 KR 2000-49344	1078687 A1 20011025 W0 2001-KR462  AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, HU, ID, IL, IN, IS, JP, KE, KG, KP, KZ, LC, LK, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, 1099583 A 20011109 KR 2000-49344  7602 A1 20020320 EP 2001-917893  AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, IE, SI, LT, LV, FI, RO 3026844 A1 20030206 US 2002-18870 KR 2000-20484 KR 2000-49344	1078687 A1 20011025 W0 2001-KR462  AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, HU, ID, IL, IN, IS, JP, KE, KG, KP, KZ, LC, LK, LR, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM  : GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, 1099583 A 20011109 KR 2000-49344  7602 A1 20020320 EP 2001-917893  AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, IE, SI, LT, LV, FI, RO 3026844 A1 20030206 US 2002-18870  PLN. INFO.: KR 2000-20484  KR 2000-49344	1078687 A1 20011025 W0 2001-KR462 20  AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, HU, ID, IL, IN, IS, JP, KE, KG, KP, KZ, LC, LK, LR, LS, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM  : GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG  1099583 A 20011109 KR 2000-49344 20  AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, IE, SI, LT, LV, FI, RO  3026844 A1 20030206 US 2002-18870 20  PLN. INFO.: KR 2000-20484 A 2000-49344 A 2000-	1078687 A1 20011025 W0 2001-KR462 200103  AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HU, ID, IL, IN, IS, JP, KE, KG, KP, KZ, LC, LK, LR, LS, LT, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM  : GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG  1099583 A 20011109 KR 2000-49344 200003  AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, IE, SI, LT, LV, FI, RO  3026844 A1 20030206 US 2002-18870 200204  PLN. INFO.: KR 2000-20484 A 2000064

AB Controlled and sustained release injectable pharmaceutical compns. for a biopharmaceutical, such as peptides and proteins are described. Processes for preparation of an injectable sustained release composition comprises (i) a step of preparing biodegradable porous microspheres having accessible ionic functional groups, (ii) a step of encapsulating a biopharmaceutical into the microspheres through ionic interaction by suspending or equilibrating the microspheres in a solution containing the biopharmaceutical, and (iii) a step of recovering and freeze-drying the biopharmaceutical-incorporated microspheres. For example, microspheres were prepared by water/oil/water double emulsion solvent evaporation method using a hydrophilic 50:50 PLGA polymer (RG 502H), which contains free carboxy end groups. Deionized water (800 mL) was added to 1 g of PLGA polymer dissolved in 2 mL of methylene chloride and emulsified by sonication for 30 s using a probe

type ultrasonic generator. This primary emulsion was dispersed into 200 mL of deionized water containing 0.5% polyvinyl alc. (weight/volume) in a vessel which connected to a constant temperature controller and mixed well by stirring for 15 min at 2500 rpm, 25° using a mixer. After mixing for another 15 min at 1500 rpm, 25°, temperature of continuous phase was increased to  $40^{\circ}$  to evaporate methylene chloride. After 1 h stirring at  $40^{\circ}$ , 1500 rpm, temperature was decreased to  $25^{\circ}$ . The hardened microspheres were collected by centrifugation and washed twice with 200 mL of deionized water, and then freeze-dried. The microspheres obtained were used for incorporation of protein drugs, i.e., ovalbumin, bovine serum albumin, human growth hormone, RNase A, or lysozyme through ionic interaction by simply soaking and equilibrating the microspheres into a buffer solution having an appropriate concentration of protein.

577-11-7, Docusate sodium

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (preparation of polymer-based injectable sustained-release microspheres for peptide and protein drugs)

577-11-7 CAPLUS RN

IT

Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, sodium salt (9CI) CN (CA INDEX NAME)

Na

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 3 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN L6

ACCESSION NUMBER:

DOCUMENT NUMBER: TITLE:

2001:434866 CAPLUS 135:37202

Compositions containing itraconazole with improved bioavailability and narrow intra- and inter-individual

variation of its absorption

INVENTOR(S):

Kwon, Jong-won; Kim, Jung-hun; Wang, Hun-sik; Jang,

Sun-woo; Bae, Woong-tak

PATENT ASSIGNEE(S):

Dong A Pharm. Co., Ltd., S. Korea

SOURCE:

PCT Int. Appl., 35 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PA.	PATENT NO.					KIND DATE			APPLICATION NO.					DATE			
WO	WO 2001041765					A1 20010614			,	WO 1999-KR854					19991231 <		
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CA	2393	737			AA		2001	0614	1	CA 1	999-	2393	737		1:	9991:	231 <
ΕP	1274	432			A1		2003	0115		EP 1	999-	9625.	55		1	9991	231

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL

JP 2003516354 T2 20030513 JP

JP 2001-543110 KR 1999-55802

19991231

PRIORITY APPLN. INFO:: KR 1999-55802 A 19991208 WO 1999-KR854 W 19991231

AB The present invention relates to compns. containing itraconazole, with both improved bioavailability, due to higher water-solubility and impressively reduced differences of pH-dependent solubility, and narrow intra- and inter-individual variation of its absorption- and a manufacturing method. formulations consist of itraconazole, a water-soluble macromol. 10-100%, solubilizer 0.1-100% and pharmaceutical acceptable additives. Itraconazole minimizes absorption variation by dozing time after food intake as well as is available for adults with hypoacidity, AIDS patients and normal people. In addition, the manufacturing method introduces the elementary process, the spray drying, thereby control of phys. properties of particles containing drug is easier. Thus, 100 g HPMC and 7 g Poloxamer were dissolved in a mixture of EtOH and CH2Cl2, and 100 g of itraconazole was added. To the resulting solution, 1 g NaCl and 1 g Mg stearate were added, and dispersed produce homogeneous spray-drying solution This solution was spray-dried at feeding rate of 150 mL/min, and atomizing pressure of 0.5 kg/cm2.

IT 577-11-7, Sodium docusate

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (compns. containing itraconazole with improved bioavailability and narrow intra- and inter-individual variation of absorption)

RN 577-11-7 CAPLUS

● Na

REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 4 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2001:380370 CAPLUS

DOCUMENT NUMBER: 135:9995

TITLE: Pharmaceuticals containing sildenafil for treating

male erectile dysfunction

INVENTOR(S): Vallabhaneni, Ramakrishna Rao

PATENT ASSIGNEE(S): Natco Pharma Ltd., India SOURCE: PCT Int. Appl., 19 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

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			SK.	ST	ጥ.ፕ.	TM.	TR.	ጥጥ.	Υ7.	IIA.	IIG.	115	117.	V/N	YII.	7.A.	7.W .	AM.	

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AB The invention relates to a novel pharmaceutical composition containing sildenafil useful for nasal administration in the treatment of male erectile dysfunction due to a variety of causes. The composition is also effective in patients with erectile dysfunction due to spinal cord injury. The pharmaceutical composition is in the form of a solution or a colloidal dispersion in a vehicle filled into a specially designed dosing device for nasal administration. The invention also provides a method for preparing the composition containing sildenafil for nasal application for the treatment of male erectile dysfunction. Thus, a formulation contained sildenafil citrate 10.000, PEG-300 30.000, glycerol 20.000, and HCl 10.000% and water to 1.0 mL.

IT 577-11-7, Docusate sodium

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (pharmaceuticals containing sildenafil for treating male erectile dysfunction)

577-11-7 CAPLUS RN

CN Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, sodium salt (9CI) (CA INDEX NAME)

## Na

CAPLUS COPYRIGHT 2006 ACS on STN ANSWER 5 OF 23

ACCESSION NUMBER:

2001:101366 CAPLUS

DOCUMENT NUMBER:

134:152659

TITLE:

Sample arrays and high-throughput testing thereof to

detect interactions

INVENTOR(S):

Putnam, David; Chen, Hongming; Galakatos, Nicholas;

Langer, Robert S.

PATENT ASSIGNEE(S):

Transform Pharmaceuticals, Inc., USA

PCT Int. Appl., 67 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

SOURCE:

English

FAMILY ACC. NUM. COUNT:

PATENT N			KINI	) I	DATE		i	APPL:	ICAT:	ION I	NO.		D	ATE	
WO 20010	09391		A1	-	2001	0208		WO 21	000-1	1520	 71 <i>7</i>		20	0000	 728 <
		G, AL,											_		. – -
	CR, CI	U, CZ,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,
	HU, I	D, IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,
	LU, L	V, MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	PL,	PT,	RO,	RU,
	SD, SI	E, SG,	SI,	SK,	SL,	TJ,	TM,	TR,	TT,	TZ,	UA,	UG,	UZ,	VN,	YU,
	ZA, ZV	W, AM,	AZ,	BY,	KG,	ΚZ,	MD,	RU,	ТJ,	TM					
RW:	GH, GI	M, KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,	CY,

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DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ,
             CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                            CA 2000-2379160
                                20010208
                                                                    20000728 <--
    CA 2379160
                          AA
                                            EP 2000-952298
    .EP 1204766
                          A1
                                20020515
                                                                    20000728
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL
                                            BR 2000-12767
    BR 2000012767
                          Α
                                20020723
                                                                    20000728
    JP 2003509657
                          Т2
                                20030311
                                            JP 2001-513646
                                                                    20000728
                                            ZA 2002-503
    ZA 2002000503
                                20030422
                                                                    20020121
                          Α
                                                                 P
                                            US 1999-146019P
                                                                    19990728
PRIORITY APPLN. INFO.:
                                            US 2000-540462
                                                                 A 20000331
                                            WO 2000-US20717
                                                                 W 20000728
```

AB The invention relates to high-throughput methods to prepare an array comprising a large number of samples, each sample consisting of a combination of components, at varying concns. and identities, and high-throughput methods to test each sample for one or more properties. Such methods allow detection or measurement of interactions or detection of lack of interactions between inactive components and active components; between multiple inactive components; or between multiple active components. The invention is particularly suited for making a large number of pharmaceutical-excipient samples at the same time, then rapidly testing each sample to detect or measure an interaction. Once such interaction is detected or measured, it can be exploited to develop optimized formulations for pharmaceutical administration. Griseofulvin formulations with enhanced solubility were identified by testing 18 excipients at different concns. and combinations.

IT 577-11-7, Sodium docusate

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(as excipients for griseofulvin formulations; sample arrays and
high-throughput testing thereof to detect interactions)

RN 577-11-7 CAPLUS

CN Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, sodium salt (9CI) (CA INDEX NAME)

Na

REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 6 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2001:31306 CAPLUS

DOCUMENT NUMBER: 134:105846

TITLE: Clear aqueous dispersions of triglycerides and

surfactants for delivery of drugs and nutrients

INVENTOR(S): Chen, Feng-Jing; Patel, Mahesh V.

PATENT ASSIGNEE(S): Lipocine, Inc., USA SOURCE: PCT Int. Appl., 103 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 13

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2001001960	A1	20010111	WO 2000-US15133	20000602 <
W: AE, AG, AL	AM, AT,	AU, AZ, BA	, BB, BG, BR, BY, CA,	CH, CN, CR,

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CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU,
           ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU,
       CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                       В1
                             20010731
                                     US 1999-345615
    US 6267985
                                                             19990630 <--
                       AA
                                        CA 2000-2375083
                                                             20000602 <--
    CA 2375083
                             20010111
                                        EP 2000-938039
                       Α1
                             20020410
                                                             20000602
    EP 1194120
           AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
           IE, SI, LT, LV, FI, RO
                       T2
                             20030128
                                        JP 2001-507455
                                                             20000602
    JP 2003503440
    NZ 516521
                       Α
                             20031128
                                        NZ 2000-516521
                                                             20000602
    AU 783077
                       B2
                             20050922
                                       AU 2000-53131
                                                             20000602
PRIORITY APPLN. INFO.:
                                        US 1999-345615
                                                          A 19990630
                                        WO 2000-US15133
                                                          W 20000602
```

The present invention relates to drug and nutrient delivery systems, and AB in particular to pharmaceutical compns. and methods for improved solubilization of triglycerides and improved delivery of therapeutic agents. Compns. of the present invention include a triglyceride and a carrier, where the carrier is formed from a combination of at least two surfactants, at least one of which is hydrophilic. Upon dilution with an aqueous solvent, the composition forms a clear, aqueous dispersion of the triglyceride and surfactants. An optional therapeutic agent can be incorporated into the composition, or can be co-administered with the composition. The invention also provides methods of enhancing triglyceride solubility and methods of treatment with therapeutic agents using these compns. Several formulations were presented of compns. that can be prepared according to the present invention using a variety of therapeutic agents. Examples of aqueous dispersions include: (1) Cremophor RH-40 0.75, Peceol 0.25, corn oil 0.40, and fenofibrate 0.10; (2) Cremophor RH-40 0.57, Crovol M-40 0.43, corn oil 0.40, and Rofecoxib 0.15; (3) Tween 80 0.70, Tween 85 0.35, Miglyol 812 0.30, Paclitaxel 0.10, and PEG 400 0.25; or (4) Kessco PEG 400 MO 0.33, corn oil 0.30, and Terbinafine 0.25 parts, resp.

IT 577-11-7, Sodium docusate

> RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (clear aqueous dispersions of triglyceride and surfactants for delivery of drugs and nutrients)

RN577-11-7 CAPLUS

Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, sodium salt (9CI) CN (CA INDEX NAME)

**N**a

REFERENCE COUNT:

2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 7 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN L6

ACCESSION NUMBER:

2000:608551 CAPLUS

DOCUMENT NUMBER:

133:213151

TITLE:

Pharmaceutical compositions and methods for improved

delivery of hydrophobic therapeutic agents

INVENTOR(S): Patel, Manesh V.; Chen, Feng-Jing

PATENT ASSIGNEE(S):

Lipocine, Inc., USA PCT Int. Appl., 98 pp.

SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE: LANGUAGE: Patent English

FAMILY ACC. NUM. COUNT: 13

PATENT INFORMATION:

KIND DATE APPLICATION NO. PATENT NO. ---------20000831 WO 2000-US165 WO 2000050007 A1 20000105 <--W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG В1 20010925 US 1999-258654 US 6294192 19990226 <--CA 2000-2365536 CA 2365536 AA20000831 20000105 <--AU 2000-22242 AU 2000022242 A5 20000105 <--20000914 AU 771659 B2 20040401 A1 20011205 EP 2000-901394 EP 1158959 20000105 <--R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO Т2 JP 2002537317 20021105 JP 2000-600619 20000105 NZ 2000-513810 NZ 513810 20040227 20000105 Α US 1999-258654 A 19990226 WO 2000-US165 W 20000105 PRIORITY APPLN. INFO.:

AB The present invention relates to triglyceride-free pharmaceutical compns. for delivery of hydrophobic therapeutic agents. Compns. of the present invention include a hydrophobic therapeutic agent and a carrier, where the carrier is formed from a combination of a hydrophilic surfactant and a hydrophobic surfactant. Upon dilution with an aqueous solvent, the composition forms a clear, aqueous dispersion of the surfactants containing the therapeutic agent. The invention also provides methods of treatment with hydrophobic therapeutic agents using these compns. A pharmaceutical composition contained cyclosporin 0.14, Cremophor RH-40 0.41, Arlacel186 0.29, sodium taurocholate 0.26, and propylene glycol 0.46 mg.

IT 577-11-7, Sodium docusate

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (pharmaceutical compns. and methods for improved delivery of hydrophobic therapeutic agents)

RN 577-11-7 CAPLUS

CN Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, sodium salt (9CI) (CA INDEX NAME)

Na

REFERENCE COUNT:

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 8 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2000:606756 CAPLUS

DOCUMENT NUMBER:

133:198661

TITLE:

Seeded microcapsules for use in tablets,

pharmaceutical agents and nutritional compounds

INVENTOR(S):

Redding, Bruce K., Jr.; Harden, Jerome

PATENT ASSIGNEE(S):

Verion Inc., USA

U.S., 14 pp., Cont. of U.S. Ser. No. 111,897. SOURCE:

CODEN: USXXAM

DOCUMENT TYPE: LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
us 6110501	Α	20000829	US 1999-226356	19990106 <
us 6149953	Α	20001121	US 1998-111897	19980708 <
PRIORITY APPLN. INFO.:			US 1993-137439 E	1 19931108
			US 1995-576636 E	1 19951221
			US 1997-908232 E	2 19970807
			US 1998-82165P F	19980417
,			US 1998-111897 A	1 19980708

AB Disclosed is a microcapsule having a core, a shell and seeds fully or partially embedded in said shell. The core and seeds are active substances which preferably function as a leavening agent. The shell is composed of either a water soluble or meltable natural polymer, including vegetable waxes. When the shell is ruptured, the active substances will react with each other and the dough mixture thereby producing a leavening effect and/or dough conditioning effect in baked goods. Seeded vitamin C microcapsules were made by mixing ascorbic acid 700 g with molten cottonseed vegetable wax 250, and microcryst. cellulose 50 g.

IT 577-11-7, Docusate Sodium

> RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (seeding material for microcapsules for use in tablets containing pharmaceutical agents and nutritional compds.)

577-11-7 CAPLUS RN

CN Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, sodium salt (9CI) (CA INDEX NAME)

REFERENCE COUNT:

14 THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 9 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2000:589894 CAPLUS

DOCUMENT NUMBER:

133:182998

TITLE:

Pharmaceutical excipient comprising microcrystalline cellulose and silica with improved compressibility

INVENTOR(S):

Staniforth, John N.; Hunter, Edward A.; Sherwood, Bob

PATENT ASSIGNEE(S):

Edward Mendell Co., Inc., USA

SOURCE:

U.S., 27 pp., Cont.-in-part of U.S. 5,866,166.

CODEN: USXXAM

DOCUMENT TYPE:

Patent English

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 6106865	Α	20000822	US 1998-37841	19980310 <
US 5585115	Α	19961217	US 1995-370576	19950109 <

	5725883			19980310			486183		19950607	<
. EP	1287823	-	.1	20030305			79378		19960105	
	R: AT, BE,									
	5866166	1		19990202			660553		19960610	
US	5725884	1		19980310			724613		19960930	
US	2001001664		.1	20010524	US	2001-	754760		20010104	<
US	6358533	I	2	20020319						
US	2002142032	1	.1	20021003	US	2001-	981319		20011016	
US	6521261	I	2	20030218					•	
US	2003099702	1	.1	20030529	US	2002-	145563		20020514	
US	6936277	I	2	20050830						
US	2003096005	1	.1	20030522	US	2003-	338361		20030108	
US	6858231	I	2	20050222						
US	2005013861	1	.1	20050120	US	2004-	850059		20040520	
US	2006008522	7	.1	20060112	US	2005-	174839		20050705	
PRIORIT	Y APPLN. INFO	.:					370576	A1	19950109	
					US	1995-	486183	A2	19950607	
					US	1996-	660553	A2	19960610	
					US	1996-	724613	A2	19960930	
					US	1996-	19546P	P	19960610	
					US	1996-	19547P	P	19960610	
					EP	1996-	903539	A3	19960718	
					US	1997-	868745	A2	19970604	
					US	1997-	992073.	A1	19971217	
					US	1998-	37841	A2	19980310	
					US	1999-	384829	B1	19990827	
					US	1999-	438646		19991112	
							754760		20010104	
					US	2001-	981319	A1	20011016	
					US	2002-	145563	<b>A</b> 1	20020514	
					US	2003-	338361	A1	20030108	

AB A composition, comprising (a) microcryst. cellulose; and (b) a compressibility augmenting agent which (i) phys. restricts the proximity of the interface between adjacent cellulose surfaces; or (ii) inhibits interactions between adjacent cellulose surfaces; or (iii) accomplishes both (i) and (ii) above, is disclosed. The composition is in the form of agglomerated particles of microcryst. cellulose and the compressibility augmenting agent in intimate association with each other. A slurry of microcryst. cellulose containing 5% silicone dioxide was spray dried to obtain a powder having an average particle size of 40-60 μm. The powder was wet granulated and wet screened through a 12 mesh screen, and dried to obtained an average particle size of 55-70 μm. Compressed tablets were prepared from the granules having good tensile strength.

IT 577-11-7, Docusate sodium

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (pharmaceutical excipient comprising microcryst. cellulose and silica with improved compressibility)

RN 577-11-7 CAPLUS

CN Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, sodium salt (9CI) (CA INDEX NAME)

Na

REFERENCE COUNT:

L6

THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

7

ACCESSION NUMBER: 2000:553455 CAPLUS

DOCUMENT NUMBER: 133:155507

TITLE: Implant comprising calcium cement and hydrophobic

liquid

INVENTOR(S):
Bohner, Marc

PATENT ASSIGNEE(S): Mathys Robert Stiftung, Switz.; Stratec Medical A.-G.

SOURCE: PCT Int. Appl., 40 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	rent 1	NO.			KINI	)	DATE		AP	PLICAT	ION I	NO.		D	ATE		
WO	20000	0458	67		A1	_	2000	0810	WO	1999-	EP68	4		1	9990	202	<
		•	•	•	JP,												
	RW:	•	•	CH,	CY,	DE,	DK,	ES,	FI, F	R, GB,	GR,	IE,	IT,	LU,	MC,	NL,	
		PT,	SE														
CA	23618	847			AA		2000	0810	CA	1999-	2361	847		1	99902	202	<
AU	99292	241			A1		2000	0825	AU	1999-	2924	1		1:	99902	202	<
AU	75491	17			B2		2002	1128									
EP	11507	722			<b>A</b> 1		2001	1107	EP	1999-	9101	83		1	9990	202	<
EP	11507	722			B1		2005	1005									
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB, I	r, LI,	LU,	NL,	SE,	MC,	PT,	IE,	FI
JP	20025	5360	75		Т2		2002	1029	JP	2000-	5969	86		1	99902	202	
AT	30580	02			E		2005	1015	AT	1999-	9101	83		1	9990	202	
US	66422	285			В1		2003	1104	US	2001-	8896	55		2	0010	719	
HK	10375	546			A1		2005	1125	HK	2001-	1079	64		2	0011	113	
PRIORITY	APPI	LN.	INFO	. :					WO	1999-	EP68	4	1	W 1	99902	202	

AB The composition comprises a hydraulic cement for implantation in the human or animal body, said hydraulic cement comprising a first component comprising a calcium source and a second component comprising water, which hardens after mixing of the components. The composition further comprises a third component with a hydrophobic liquid The composition allows to obtain a cement with open macroporosity enabling a rapid bone ingrowth. A mixture of α-tri-calcium phosphate 8, precipitated tricalcium phosphate 0.8, calcium cement 0.5 g, Cremophor EL 0.001, and Tegosoft M 8.0 mL were stirred for 4 min. The mixture was then poured into a syringe and injected into a cavity. After hardening, the cavity was filled with an open macroporous calcium phosphate structure.

IT 577-11-7, Docusate sodium

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (implant comprising calcium cement and hydrophobic liquid)

RN 577-11-7 CAPLUS

CN Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, sodium salt (9CI) (CA INDEX NAME)

Na

REFERENCE COUNT: 14 THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 11 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2000:290723 CAPLUS

DOCUMENT NUMBER: 132:307237

TITLE: A trypsinized and Coomassie Brilliant Blue-stained

Leishmania promastigote composition useful for the early diagnosis of visceral leishmaniasis and a

process for preparing the same

INVENTOR(S):

Girish, Kumar Jain; Suman, Tiwari; Suman, Gupta;

Katiyar, Jagdish Chandra

PATENT ASSIGNEE(S):

Council of Scientific and Industrial Research, India

SOURCE:

Eur. Pat. Appl., 13 pp.

CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAI	CENT	NO.			KIN	D	DATE			APPL	ICAT	ION	NO.		D	ATE	•	
	<b>-</b>			•		-									_			
ΕP	9977	34			A1		2000	0503		EP 1	998-	8903	17		1	9981	029	<
	ъ.	ΔΨ	BF	CH	DE	DΚ	FC	FD	CB	CP	TΨ	T.T	T.T	MT.	SF	MC	рŢ	

AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,

IE, SI, LT, LV, FI, RO

PRIORITY APPLN. INFO.:

EP 1998-890317

19981029

A composition for the early diagnosis of visceral leishmaniasis comprises trypsinized and Coomassie Brilliant Blue-stained Leishmania promastigotes and a protein stabilizing solute in the ratios of 5 million: 0.0001 mg to 100 million: 1.00 mg. The protein stabilizing solute is surfactant, glycerol, sucrose, etc. The composition is used to test serum samples by direct agglutination test.

128-49-4, Docusate calcium 577-11-7, IT

Docusate sodium 7491-09-0, Docusate potassium

RL: ARU (Analytical role, unclassified); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses)

(as protein-stabilizing solute; trypsinized and Coomassie Brilliant Blue-stained Leishmania promastigote composition useful for early diagnosis of visceral leishmaniasis and its preparation)

128-49-4 CAPLUS RN

Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, calcium salt (9CI) CN (CA INDEX NAME)

### ●1/2 Ca

RN 577-11-7 CAPLUS

Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, sodium salt (9CI) CN (CA INDEX NAME)

## Na

RN 7491-09-0 CAPLUS

Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, potassium salt CN (9CI) (CA INDEX NAME)

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THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 12 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN L6

ACCESSION NUMBER: 2000:123188 CAPLUS

DOCUMENT NUMBER:

132:171126

Flocculated suspension of megestrol acetate TITLE:

Ragunathan, Narayan; Chao, James C.; Femia, Robert A.; INVENTOR(S):

Ross, Malcolm S. F.

PATENT ASSIGNEE(S): Pharmaceutical Resources, Inc., USA

U.S., 5 pp. CODEN: USXXAM SOURCE:

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION: DAMENIM NO

PA'	TENT	NO.			KIN		DATE		i	APPL	ICAT	ION 1	NO.		I	ATE		
	6028				Α								_			.9980		
WO	2001															9991		
	W:	AL,	AM,	AT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CU,	CZ,	DE,	
		DK,	EE,	ES,	FI,	GB,	GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	
		KG,	KP,	KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	MN,	MW,	
		MX,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TR,	
		TT.	UA.	UG.	UZ.	VN.	YU,	ZA.	ZW.	AM.	AZ.	BY.	KG.	KZ.	MD.	RU,	TJ.	TM
	RW:		•	•	•		•			•		•			•	CY,		
			-										-	-		BJ,		
			•					ML,	•	•			-	,	J.,	20,	·-,	
זומ	9962	•	•	•	•	•	•	•	•	•	•	•			1	.9991	007	<b>&lt;</b>
	6268							0731								.9991		
	2001									JS 2	001-	1512	ρŢ		-	20010	109	<
	6593				B2			0715										
US	2002	1734					2002	1121	1	JS 2	002-	1368	23		2	20020	430	
US	6593	320			B2		2003	0715										
PRIORIT	Y APP	LN.	INFO	. :					1	JS 1	998-	6324	1.		A 1	9980	420	
									1	WO 1	999-	US23	340		A 1	9991	007	
																9991		
																20010		
. מ כומ	1		1		1		~~~											

AB A novel oral antineoplastic composition comprises a stable flocculated suspension in water containing megestrol acetate, ≥1 of PEG, propylene glycol, glycerol, and sorbitol, and a surfactant, provided polysorbate and PEG are not simultaneously present. Any surfactant, regardless of the length of the hydrophobic contact area on its hydrophobic group, can effectively wet megestrol acetate and form a stable flocculated suspension, provided ≥1 of the other named compds. is present. Thus, a suspension was prepared containing megestrol acetate 4.000, glycerol 5.000, sorbitol 15.000, docusate Na (surfactant) 0.002, xanthan gum 0.250, NaOBz 0.200, citric acid 0.300, Na citrate 0.060, sucrose 5.000, lemon flavoring 0.080, and H2O 70.108 weight%.

ΙT 577-11-7, Docusate sodium

> RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (surfactant; flocculated suspension of megestrol acetate)

RN 577-11-7 CAPLUS

#### Na

IT 5138-18-1D, Sulfosuccinic acid, esters with fatty alcs.
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(surfactants; flocculated suspension of megestrol acetate)

RN 5138-18-1 CAPLUS

CN Butanedioic acid, sulfo- (9CI) (CA INDEX NAME)

 $^{\rm SO3H}$   $^{\rm HO_2C-CH-CH_2-CO_2H}$ 

REFERENCE COUNT: 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 13 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1999:763873 CAPLUS

DOCUMENT NUMBER:

132:15626

TITLE:

Preparation of efavirenz and compressed tablet

containing efavirenz

INVENTOR(S):

Batra, Udit; Higgins, Raymond J.; Thompson, Karen C.;

Katdare, Ashok V.

PATENT ASSIGNEE(S):

SOURCE:

Merck and Co., Inc., USA PCT Int. Appl., 31 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PA	CENT															ATE		
WO	9961															9990!	524	<
	W:	ΑE,	AL,	AM,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	CA,	CN,	CU,	CZ,	EE,	GD,	
								IS,										
		MD,	MG,	MK,	MN,	MX,	NO,	NZ,	PL,	RO,	RU,	SG,	SI,	SK,	SL,	TJ,	TM,	
		TR,	TT,	UA,	US,	UZ,	VN,	YU,	ZA,	AM,	AZ,	BY,	KG,	KZ,	MD,	RU,	TJ,	TM
	RW:	GH,	GM,	KE,	LS,	MW,	SD,	SL,	SZ,	UG,	ZW,	AT,	BE,	CH,	CY,	DE,	DK,	
		ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,	CG,	
		CI,	CM,	GA,	GN,	GW,	ML,	MR,	NE,	SN,	TD,	TG						
US	2001	0143	52		A1		2001	0816		US 1	999-	3126	17		1:	9990!	517	<
CA	2332	876			AA		1999	1202		CA 1	999-	2332	876		1	9990!	524	<
ΑU	9942	010			A1		1999	1213		AU 1	999-	4201	0		1	9990!	524	<
ΑU	7611	82			B2		2003	0529										
EΡ	1083	901			A1		2001	0321		EP 1	999-	9257	93		1	9990!	524	<
EP	1083	901			B1		2003	0416										
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	PT,	ΙE,	
		SI,	LT,	LV,	FI,	RO												
JΡ	2002	5162	81		Т2		2002	0604		JP 2	000-	5504	86		1	9990!	524	
AT	2373	32			E		2003	0515		AT 1	999-	9257	93		1:	9990!	524	
ΕP	1332	757			<b>A</b> 1		2003	0806		EP 2	003-	7605	4		1	9990!	524	
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	PT,	IE,	

A 50 % drug loaded compressed tablet formulation for efavirenz (I) is AB disclosed. I is a non-nucleoside reverse transcriptase inhibitor being studied clin. for use in the treatment of HIV infections and AIDS. I was prepared by grignard cyclization of 4-chloro-2-(trifluoroacetyl)aniline. Tablets containing 50% I were prepared The core were comprised I 950, microcryst. cellulose 380, hydroxypropyl cellulose 60.8, croscarmellose sodium 95, sodium lauryl sulfate 19 g, lactose hydrous spray dried 19.8, magnesium stearate 1% and water 1.045 L; and the film coating material comprised hydroxypropyl cellulose 8.54, hydroxypropyl Me cellulose 8.54, titanium dioxide 3.42 mg, and water 94%.

WO 1999-US11464

W 19990524

577-11-7, Docusate sodium ΙT

> RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (preparation of efavirenz and compressed tablet containing efavirenz) 577-11-7 CAPLUS

RN

Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, sodium salt (9CI) CN (CA INDEX NAME)

Na

REFERENCE COUNT: THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS 3 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 14 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1999:27714 CAPLUS

DOCUMENT NUMBER:

TITLE: Pharmaceutical compositions containing synergistic

acetaminophen and cisapride

Hough, Douglas R.; Nelson, Edward B.; Raffa, Robert B. INVENTOR(S):

PATENT ASSIGNEE(S): McNeil-PPC, USA

SOURCE: PCT Int. Appl., 16 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

English LANGUAGE:

FAMILY ACC. NUM. COUNT:

	PAT	CENT 1	10.			KINI	)	DATE		į	APPL:	ICAT:	ION 1	10.		DA	ATE	
	WO	98586	547			A1	•	 1998:	1230	,	WO 19	 997-τ	JS108	358		19	9700	523 <
		W:	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CU,	CZ,	DE,
			DK,	EE,	ES,	FI,	GB,	GE,	HU,	IL,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,
			LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,
			RO,	RU,	SD,	SE,	SG,	SI,	SK,	TJ,	TM,	TR,	TT,	UA,	UG,	UZ,	VN,	AM,
			ΑZ,	BY,	KG,	ΚZ,	MD,	RU,	TJ,	TM								
		RW:	GH,	KE,	LS,	MW,	SD,	SZ,	ŪG,	ZW,	ΑT,	BE,	CH,	DE,	DK,	ES,	FI,	FR,
			GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,
			GN,	ML,	MR,	NE,	SN,	TD,	TG									
	ΑU	97432	257			<b>A1</b>		19990	0104		AU 19	997-	4325	7		19	9970	523 <
PRIOF	RITY	APPI	LN.	INFO	. :					1	WO 19	997 <b>-</b> 1	US10	358	7	A 19	9706	523
AB	Dis	sclose	ed a	re co	ompns	s. co	ompr	ising	g ace	etam	inopl	hen	(I) a	and o	cisap	pride	e (I:	[) and

methods for their use in analgesia. When acetaminophen and cisapride are administered in combination, their analgesic pharmacol. effects are superadditive. A mixture of 30 mg I and 30 mg II was orally administered to mice followed by injection of 5.5 mg/kg acetylcholine bromide 30 min later. The ED50 of I and II decreased from 169.5 and 34.6 mg to 9.1 for each, resp. and 13 out of 15 mice showed no writhing.

IT 577-11-7, Docusate sodium

> RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (pharmaceutical compns. containing synergistic acetaminophen and cisapride)

577-11-7 CAPLUS RN

Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, sodium salt (9CI) CN (CA INDEX NAME)

Na

REFERENCE COUNT:

2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 15 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1999:27707 CAPLUS

DOCUMENT NUMBER:

130:86181

TITLE:

Pharmaceutical formulations containing ibuprofen and

diphenhydramine analgesics

INVENTOR(S):

Hough, Douglas R.; Nelson, Edward B.; Raffa, Robert B.

PATENT ASSIGNEE(S):

McNeil-PPC, USA

SOURCE:

PCT Int. Appl., 15 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

	PATENT	NO.			KIN	D :	DATE		•				NO.		D	ATE	
	WO 985	8640			A1		1998	1230	,						19	9970	623 <
	W:	AL,	AM,	AT,	ΑU,	AZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CU,	CZ,	DE,
		DK,	EE,	ES,	FI,	GB,	GE,	HU,	IL,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,
		LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,
							SI,										
		AZ,	BY,	KG,	KZ,	MD,	RU,	TJ,	TM							-	•
	RV	: GH,	KE,	LS,	MW,	SD,	SZ,	UG,	ZW,	AT,	BE,	CH,	DE,	DK,	ES,	FI,	FR,
		GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	BJ,	CF,	CG,	CI,	CM,	GA,
		GN.	ML,	MR.	NE,	SN,	TD,	TG	•	•	•	•	•	-	•	•	•
	AU 974	3256	•	•	A1	·	1999	0104		AU 19	997-	4325	6		19	9970	623 <
PRIO	RITY AF	PLN.	INFO	. :					,	WO 19	997-1	US10	857		A 19	9970	623
AB	Disclo																
	and me																•
	dipher							_				-				ects	are
	supera	_										-					
	_															_	tested
	in ace		-	_		_											
	writhi	-						_				0			_, 0,		-
IT	577-11	_	0016	ata	sodii	ım											
	3,,-11	. ,, ,		acc	J.J.				_								

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (pharmaceutical formulations containing ibuprofen and diphenhydramine analgesics)

577-11-7 CAPLUS RN

Na

REFERENCE COUNT:

3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 16 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1998:226811 CAPLUS

DOCUMENT NUMBER:

128:286378

TITLE:

Synergistic analgesic combination containing

acetaminophen and dimenhydrinate

INVENTOR(S):

Hough, Douglas R.; Nelson, Edward B.; Raffa, Robert B.

PATENT ASSIGNEE(S):

McNeil-PPC, Inc., USA

SOURCE:

U.S., 5 pp.

DOCUMENT TYPE:

CODEN: USXXAM Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

	PA	FENT	NO.			KIN	D	DATE					ION :			D	ATE	
		5739 9858									us 1	996-	6670	54				620 < 623 <
		W:							BB,									
			DK,	EE,	ES,	FI,	GB,	GE,	HU,	IL,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,
			LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,
									SK,									
									TJ.		•	•	•	•	•	•		•
		RW:	GH.	KE.	LS.	MW.	SD.	SZ.	ΰĠ,	ZW.	AT.	BE.	CH.	DE.	DK.	ES.	FI.	FR.
			•	•	•	•			NL,		•	•	•	•	•			•
								TD,		,		,	,	,	•			
	WO	9858	•	•	•	•	•	•		,	WO 1	997-	us10	918		1	9970	623 <
								-	BB,									
		***	•		•	•			HU,				•	•				-
			•		•			•	MD,	•	•	•	•	•	-			•
									SK,									
			•	•	•	•	•	•	TJ,	•	,	111,	,	011,	00,	02,	•117	,
		ъw.	•	•	•	•		•	UG,		ΔT	BF	СН	DF	DΚ	FS	FT	FD
		1100 .	-		-			-	NL,	•	-			•	-		-	-
								TD,		E 1,	JE,	Dr,	БО,	CL,	CG,	CI,	CI1,	GA,
	7.11	9735	•	ти,	•	•	•	•			זות 1	007_	2500	1		1	9970	623 <
		9742												_				623 <
DDTC						ΑI		1333	0104							A 1		
PRIC	KII.	Y APP	ти.	INFO	• •						US 1							
											WO 1					A 1		
		,									WO 1					A 1		
AB	Di	scros	ed a	re c	ompn.	s. c	ompr	ısin	q ac	etam	ınop.	nen	(T)	and 4	alme:	nnyd.	rına	te (II)

Disclosed are compns. comprising acetaminophen (I) and dimenhydrinate (II) and methods for their use in analgesia. When acetaminophen and dimenhydrinate are within certain ratios, their pharmacol. effects are superadditive. Thus, 600 mg tablets containing 500 mg I and 5 mg II were prepared The synergistic activity of combination of I:II (10:1) was shown in mouse acetylcholine bromide-induced abdominal constriction assay.

ΙT 577-11-7, Docusate sodium

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (synergistic analgesic combination containing acetaminophen and

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dimenhydrinate)
N 577-11-7 CAPLUS
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RN 577-11-7 CAPLUS ĆN Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, sodium salt (9CI) (CA INDEX NAME)

Na

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ALCOND. ILLE CITATIONS INTERESTED IN THE ALCOND.

L6 ANSWER 17 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1998:25161 CAPLUS

DOCUMENT NUMBER: 128:106421

TITLE: Synergistic analgesics comprising acetaminophen and

meclizine

INVENTOR(S): Hough, Douglas R.; Nelson, Edward B.; Raffa, Robert B.

PATENT ASSIGNEE(S): McNeil-Ppc, USA

SOURCE: PCT Int. Appl., 19 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PENT				KIN	D	DATE		1	APPL	ICAT	ION I	NO.		D	ATE	
	9748				A1	_	1997	1224	Ī	WO 1:	997-	us10	922		1:	9970	620 <
	W:	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CU,	CZ,	DE,
		DK,	EE,	ES,	FI,	GB,	GE,	HU,	IL,	IS,	JP,	KE,	KG,	KP,	KR,	ΚZ,	LC,
		LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,
		RO,	RU,	SD,	SE,	SG,	SI,	SK,	ТJ,	TM,	TR,	TT,	UA,	UG,	UZ,	VN,	AM,
		AZ,	BY,	KG,	ΚZ,	MD,	RU,	ТJ,	TM								
	RW:	GH,	KE,	LS,	MW,	SD,	SZ,	UG,	ZW,	AT,	BE,	CH,	DE,	DK,	ES,	FI,	FR,
•		GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,
		GN,	ML,	MR,	NE,	SN,	TD,	TG									
ZA	9705	444			Α		1998	1221	:	ZA 1:	997-	5444			19	9970	619 <
AU	9735	763			A1		1998	0107	1	AU 1	997-	3576	3		1	9970	620 <
PRIORIT	Y APP	LN.	INFO	.:					Ī	US 1	996-	6678	34	1	A 1	9960	620
									Ī	WO 1:	997-	US10	922	1.	W 1	9970	620

AB Disclosed are synergistic compns. comprising acetaminophen (I) and meclizine (II) for use as analysesics. The ED50 of I and II in mouse acetylcholine bromide induced abdominal constriction assay was 169.5 and 159.7 mg/kg orally. The ED50 of combination of I and II (1:10) was 1.6 and 16.1 mg/kg orally.

IT 577-11-7, Docusate sodium

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (synergistic analgesics comprising acetaminophen and meclizine)

RN 577-11-7 CAPLUS

CN Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, sodium salt (9CI) (CA INDEX NAME)

Na

L6 ANSWER 18 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1996:58346 CAPLUS

DOCUMENT NUMBER: 124:97804

TITLE: Agglomerated hydrophilic complexes with multi-phasic

release characteristics

INVENTOR(S): Baichwal, Anand R.; Staniforth, John N.

PATENT ASSIGNEE(S): Edward Mendell Co., Inc., USA

SOURCE: U.S., 23 pp. Cont.-in-part of U.S. 922,312.

CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
		<b></b>		
US 5478574	Α	19951226	US 1993-94504	19930720 <
US 5472711	Α	19951205	US 1992-922312	19920730 <
IL 106253	<b>A1</b>	19980310	IL 1993-106253	19930706 <
AU 9341855	<b>A</b> 1	19940203	AU 1993-41855	19930708 <
AU 669531	B2	19960613		
HU 67622	A2	19950428	HU 1993-2104	19930721 <
CA 2101189	AA	19940131	CA 1993-2101189	19930723 <
CA 2101189	С	19990921		
JP 06172221	A2	19940621	JP 1993-204461	19930728 <
US 5670168	Α	19970923	US 1996-664792	19960617 <
PRIORITY APPLN. INFO.:			US 1992-922312	A2 19920730
			US 1995-467583	B1 19950606

AB The present invention relates to a controlled release formulation which includes a therapeutically active medicament, a heterodisperse gum matrix, a pharmaceutically acceptable diluent, and an effective amount of a pharmaceutically acceptable surfactant and/or wetting agent to provide a multi-phasic controlled release of a therapeutically active medicament. An excipient granulation is prepared containing xanthan and locust bean gums.

IT 577-11-7, Docusate sodium

RL: MOA (Modifier or additive use); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(agglomerated hydrophilic complexes with multi-phasic controlled-release characteristics)

RN 577-11-7 CAPLUS

CN Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, sodium salt (9CI) (CA INDEX NAME)

L6 ANSWER 19 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1995:795352 CAPLUS

DOCUMENT NUMBER: 123:208884

TITLE: Liquid polymer compositions for sustained drug release

INVENTOR(S): Friedman, Michael; Sintov, Amnon PATENT ASSIGNEE(S): Perio Products, Ltd., Israel

SOURCE: U.S., 53 pp. Cont.-in-part of U.S. 5,330,746.

CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 6

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
PATENT NO.  US 5438076 US 5330746 US 5139768 US 5403577 US 5849266 US 5639795 US 5648399 PRIORITY APPLN. INFO.:	KIND A A A A A A A	DATE 19950801 19940719 19920818 19950404 19981215 19970617 19970715	APPLICATION NO	B2	DATE 19930104 < 19890621 < 19910228 < 19920612 < 19950404 < 19950425 < 19950425 < 19880503 19890131 19890621
			US 1990-532328 US 1991-662985 US 1992-898096 US 1993-2481	A1 A1	19900605 19910228 19920612 19930104

AB The treatment of gingivitis, oral plaque and oral or dermatol. fungal infections comprises of administration of a liquid methacrylic acid copolymer composition that contains a release-adjusting agent and a pharmacol. agent. The composition forms a solid film upon drying, and is capable of accomplishing the sustained release of the pharmacol. agent such as to permit its use in the treatment or prevention of dental or dermatol. conditions. A liquid polymer composition containing lysine 0.3, Eudragit L 54.7, cetyl pyridinium chloride (CPC) 30, and PEG 400 15%, resp., was prepared and dried, and the cumulative release of CPC from the film produced by drying was observed

IT 577-11-7, Sodium docusate

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(liquid polymer compns. for sustained drug release)

RN 577-11-7 CAPLUS

CN Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, sodium salt (9CI) (CA INDEX NAME)

Na

L6 ANSWER 20 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1994:144201 CAPLUS

DOCUMENT NUMBER: 120:144201

TITLE: Oral dosage forms containing agglomerated hydrophilic

complexes with multi-phasic release characteristics

INVENTOR(S): Baichwal, Anand R.; Staniforth, John N.

PATENT ASSIGNEE(S): Edward Mendell Co., Inc., USA

Eur. Pat. Appl., 25 pp. SOURCE:

CODEN: EPXXDW

Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

DOCUMENT TYPE:

PATENT NO.				KIND		DATE		APPLICATION NO.				DATE				
EP	581676			A2		199402	02	EP	1993-	40195	57		1	9930'	727	<
EP	581676 R: AT,	BF		B1 DE	DΚ	200512		GB GI	D TE	TT	T.T	T.II	мс	NT.	ייים	<b>SF</b>
	5472711	DE,	CII,	Α		199512	05	บร	1992-	92231	L2		1	9920	730	<
	106253			A1		199803			1993-							
	9341855 669531			A1 B2		199402 199606		AU	1993-	41855	)		1	9930	708	<
	67622			A2		199504							_	9930	721	<
	2101189 2101189			AA C		199401 199909		CA	1993-	21011	.89		1	9930	723	<
	1582205			A2		200510		EP	2005-	10605	66		1	9930.	727	
	R: AT,	-					-			-	-	-	-	-	-	
	06172221					199406								9930'	728	<
US	5670168			Α		199709	23	US	1996-	66479	92		1	9960	617	<
PRIORITY APPLN. INFO.:							US	1992-	92231	.2	1	A 1:	9920	730		
								EP	1993-	40195	57	1	A3 1	9930	727	
								US	1995-	46758	33	]	B1 1	9950	606	

AB A controlled-release oral formulation includes a therapeutically active medicament, a heterodisperse gum matrix, a pharmaceutically acceptable diluent, and an effective amount of a surfactant and/or wetting agent to provide a multi-phasic controlled release of a therapeutically active medicament. For example, a controlled-release excipient for multiphasic dosage forms contained xanthan gum 25.0, locust bean gum 25.0, Na lauryl sulfate 5.0, and dextrose 45.0%.

IT 577-11-7, Docusate sodium

RL: BIOL (Biological study)

(controlled-release oral pharmaceuticals containing, as wetting agent)

RN577-11-7 CAPLUS

CN Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, sodium salt (9CI) (CA INDEX NAME)

# Na

ANSWER 21 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1993:577482 CAPLUS

DOCUMENT NUMBER:

119:177482

TITLE:

SOURCE:

The inhibitory effect of spermicidal agents on

replication of HSV-2 and HIV-1 in vitro

AUTHOR(S):

Jennings, R.; Clegg, A.

CORPORATE SOURCE:

Med. Sch., Univ. Sheffield, Sheffield, S10 2RX, UK

Journal of Antimicrobial Chemotherapy (1993

), 32(1), 71-82

CODEN: JACHDX; ISSN: 0305-7453

DOCUMENT TYPE:

Journal

LANGUAGE:

English

Five spermicides, including nonoxynol-9, were assessed under in vitro conditions for their inhibitory activity against two viruses capable of spread by sexual intercourse: herpes simplex virus type 2 (HSV-2) and the human immunodeficiency virus type 1. A further eight com. available spermicidal prepns. containing varying concns. of either nonoxynol-9 or nonoxynol-11 were also assessed for activity against HSV-2. All spermicides and spermicidal prepns. tested showed inhibitory activity against both viruses over periods of time ranging from 30 s to 5 min. This activity was dependent on the concentration of spermicide to which the viruses were exposed.

IT 577-11-7, Sodium docusate

RL: BIOL (Biological study)

(herpes simplex 2 and HIV-1 viruses inhibition by)

RN 577-11-7 CAPLUS

CN Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, sodium salt (9CI) (CA INDEX NAME)

#### Na

L6 ANSWER 22 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1993:455828 CAPLUS

DOCUMENT NUMBER: 119:55828

TITLE: Status of certain additional over-the-counter drug

category II and III active ingredients

CORPORATE SOURCE: United States Food and Drug Administration, Rockville,

MD, 20857, USA

SOURCE: Federal Register (1993), 58(88), 27636-44,

10 May 1993

CODEN: FEREAC; ISSN: 0097-6326

DOCUMENT TYPE: Journal LANGUAGE: English

AB Certain over-the-counter drugs are not generally recognized as safe and effective or are misbranded under the Federal Food, Drug, and Cosmetic Act. The list includes digestive, external analgesic, insect bite and sting, poison ivy, skin protectant, diaper rash, topical antifungal, and oral analgesic products.

IT 577-11-7, Docusate sodium

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (over-the-counter prepns. containing, stds. for)

RN 577-11-7 CAPLUS

CN Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, sodium salt (9CI) (CA INDEX NAME)

### Na

L6 ANSWER 23 OF 23 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1993:161252 CAPLUS

DOCUMENT NUMBER: 118:161252

TITLE: Quantification of the in vitro activity of some

compounds with spermicidal activity

AUTHOR(S): Chantler, Eric; Fisher, Helen; Solanki, Suren;

Elstein, Max

CORPORATE SOURCE: Dep. Obstetr. Gynaecol., Univ. Hosp. South Manchester,

JK

SOURCE: Contraception (1992), 46(6), 527-36

CODEN: CCPTAY; ISSN: 0010-7824

DOCUMENT TYPE: Journal LANGUAGE: English

AB The in vitro spermicidal activity of the commonly used surfactant spermicides and the antiseptic chlorhexidine, were quantified in a statistically reproducible manner, using donor semen and image capture anal. The spermicidal activity was expressed as the Ed50 under defined assay conditions. Using these parameters, the order of spermicidal activity was: Menfegol > nonoxynol-9 ≈ benzalkonium chloride > sodium docusate > chlorhexidine. These differences were statistically significant.

IT 577-11-7, Sodium docusate

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study)

(spermicidal activity of)

RN 577-11-7 CAPLUS

CN Butanedioic acid, sulfo-, 1,4-bis(2-ethylhexyl) ester, sodium salt (9CI) (CA INDEX NAME)

Na

eview at:

http://www.cas.org/infopolicy.html

Uploading C:\Program Files\Stnexp\Queries\438b.str

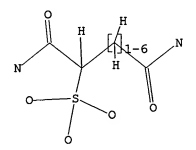
STRUCTURE UPLOADED L1

=> d

L1 HAS NO ANSWERS

L1

STR



Structure attributes must be viewed using STN Express query preparation.

### => s 11

# REG1stRY INITIATED

Substance data SEARCH and crossover from CAS REGISTRY in progress... Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

SAMPLE SEARCH INITIATED 15:01:47 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED -45 TO ITERATE

100.0% PROCESSED

45 ITERATIONS

3 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS:

ONLINE \*\*COMPLETE\*\*

BATCH

\*\*COMPLETE\*\* 1302

PROJECTED ITERATIONS:

498 TO

163

PROJECTED ANSWERS:

3 TO

L2

3 SEA SSS SAM L1

L3 3 L2

=> s 13 and py<2002 21808290 PY<2002

L41 L3 AND PY<2002

=> d ibib abs hitstr

ANSWER 1 OF 1 CAPLUS COPYRIGHT 2006 ACS on STN L4

ACCESSION NUMBER:

1985:160051 CAPLUS

DOCUMENT NUMBER:

102:160051

TITLE:

Preparation of surfactants with demonstrated

pharmacological activity

AUTHOR(S):

Kabachnyi, V. I.; Chernykh, V. P.; Kabachnyi, G. I.;

Sopel'nik, E. M.

CORPORATE SOURCE:

Khar'k. Farm. Inst., Kharkov, USSR

SOURCE:

Khimiko-Farmatsevticheskii Zhurnal (1985),

19(1), 43-6

CODEN: KHFZAN; ISSN: 0023-1134

DOCUMENT TYPE:

Journal

LANGUAGE:

Russian

OTHER SOURCE(S):

CASREACT 102:160051

Sixteen surfactant sulfosuccinic acid heterylamides were prepared and tested for pharmacol. activity and toxicity in mice. Several of the compds. exhibited anti-inflammatory activity comparable to that of butadione, and several caused lowering of blood sugar levels comparable to those produced by butamide.

IT 95896-27-8P

> RL: BAC (Biological activity or effector, except adverse); BPR (Biological process); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); PROC (Process); USES (Uses)

(preparation and pharmacol. of)

RN 95896-27-8 CAPLUS

CN 2-Butanesulfonic acid, 1-(methylamino)-4-[[5-(2-methylpropyl)-1,3,4thiadiazol-2-yl]amino]-1,4-dioxo-, monosodium salt (9CI) (CA INDEX NAME)

Na

=> s 11 full

## REG1stRY INITIATED

Substance data SEARCH and crossover from CAS REGISTRY in progress... Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

FULL SEARCH INITIATED 15:02:42 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED -849 TO ITERATE

100.0% PROCESSED 849 ITERATIONS 71 ANSWERS

SEARCH TIME: 00.00.01

L5 71 SEA SSS FUL L1

L6 26 L5

=> s 16 and py<2002 21808290 PY<2002

19 L6 AND PY<2002 1.7

=> d 1-19 ibib abs hitstr

ANSWER 1 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1997:672874 CAPLUS

DOCUMENT NUMBER: 127:294130 TITLE:

Redispersible powders based on carboxylated

butadiene-styrene and/or -acrylonitrile copolymers Rothenhaeuser, Bernd; Kiesel, Volker; Kuehn, Hartmut;

Elsaesser, Dominik

PATENT ASSIGNEE(S):

Buna Sow Leuna Olefinverbund Gmbh, Germany

SOURCE:

Ger. Offen., 7 pp. CODEN: GWXXBX

DOCUMENT TYPE:

INVENTOR(S):

Patent German

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA'	TENT NO.			KINI		APPLICATION NO.		DATE
DE	19613302	2				DE 1996-19613302		19960403 <
DE	19710380	)		<b>A</b> 1	19980917	DE 1997-19710380		19970313 <
WO	9738042			<b>A</b> 1	19971016	WO 1997-DE607		19970325 <
	W: AU,	BR,	CA,	CN,	CZ, JP, KR,	MX, PL, RU, TR, US		
	RW: AT,	BE,	CH,	DE,	DK, ES, FI,	FR, GB, GR, IE, IT,	LU,	MC, NL, PT, SE
AU	9728853			A1	19971029	AU 1997-28853		19970325 <
EP	891389			A1	19990120	EP 1997-922815		19970325 <
EP	891389			B1	20020724			
-	R: AT,	BE,	CH,	DE,	ES, FR, GB,	IT, LI, NL, SE, FI		
JP	11508959	•		T2	19990803	JP 1997-535734		19970325 <
JP	3222473			В2	20011029			
RU	2178427			C2	20020120	RU 1998-120109		19970325
AT	221095			E	20020815	AT 1997-922815		19970325
US	20021200	043		A1	20020829	US 1998-155306		19980924
PRIORIT	Y APPLN.	INFO	.:			DE 1996-19613302	7	A 19960403
						DE 1997-19710380	1	A 19970313
						WO 1997-DE607	V	v 19970325

Free-flowing, lump-free powders for curable films with good tensile strength/elongation balance, useful in construction work, are manufactured by spray-drying of carboxylated butadiene-styrene latexes with ≥1 of alkylated disulfophenyl ether salts, caseinates and Nalkylsulfosuccinamide salts as spray-drying aids.

IT 116453-32-8D, 2-Butanesulfonic acid, 1,4-diamino-1,4-dioxo-,

N-alkyl derivs., salts

RL: MOA (Modifier or additive use); USES (Uses)

(spray-drying aids; manufacture of redispersible powders from carboxylated butadiene-styrene and/or -acrylonitrile copolymer latexes)

RN 116453-32-8 CAPLUS

CN 2-Butanesulfonic acid, 1,4-diamino-1,4-dioxo- (9CI) (CA INDEX NAME)

ANSWER 2 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1995:878829 CAPLUS

DOCUMENT NUMBER:

123:290451

TITLE:

SOURCE:

Amides of sulfosuccinic acid and polyhydroxyalkylamine

for use as surfactants

INVENTOR(S):

Fabry, Bernd

PATENT ASSIGNEE(S):

Henkel K.-G.a.A., Germany

Ger. Offen., 11 pp.

CODEN: GWXXBX

DOCUMENT TYPE:

Patent

LANGUAGE:

German

\_\_\_\_

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.

KIND DATE APPLICATION NO.

DATE

DE 4336802 A1 19950511 DE 1993-4336802 19931028 <-PRIORITY APPLN. INFO.: DE 1993-4336802 19931028

AB The title amides, e.g., mono- and diamides prepared by amidation of 1 mol maleic anhydride with 1 or 2 mol N-methylglucamine followed by sulfonation of the double bond of the maleic residue, show good foaming properties and skin compatibility and are useful in detergent compns. for dishwashing and laundering, etc.

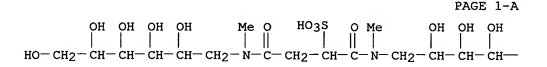
IT 169318-67-6P

RL: IMF (Industrial manufacture); PRP (Properties); TEM (Technical or engineered material use); PREP (Preparation); USES (Uses)

(surfactants; preparation and use in foaming detergent compns. with mildness to skin)

RN 169318-67-6 CAPLUS

CN D-Glucitol, 1,1'-[(1,4-dioxo-2-sulfo-1,4-butanediyl)bis(methylimino)]bis[1-deoxy-, monoammonium salt (9CI) (CA INDEX NAME)



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PAGE 1-B

L7 ANSWER 3 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1995:78033 CAPLUS 122:58115

DOCUMENT NUMBER: TITLE:

Cords of continuous filaments based on polyamides and

their manufacture Cavalie, Charles

INVENTOR(S):
PATENT ASSIGNEE(S):

Rhone-Poulenc Fibres, Fr.

SOURCE:

Fr. Demande, 11 pp.

CODEN: FRXXBL

DOCUMENT TYPE:

Patent

LANGUAGE:

French

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
FR 2692600	<b>A</b> 1	19931224	FR 1992-7688	19920618 <
FR 2692600	B1	19940826		
PRIORITY APPLN. INFO.:			FR 1992-7688	19920618

AB Title cords, especially nylon 66, have a total titer of >110 dtex, moisture content >15%, and size content 0.05-0.20% based on the weight of the cord. The process allows the manufacture of large titer cords without braiding and provides good adhesion between filaments and good cutting properties to form short fibers. The size may be a fatty amide sulfite and the short fibers may be used in electrostatic projection.

IT 94200-33-6, Sopromine 1686

RL: USES (Uses)

(sizes, for large titer polyamide fiber cords)

RN 94200-33-6 CAPLUS

CN Octadecanoic acid, (1,4-dioxo-2-sulfo-1,4-butanediyl)bis(imino-2,1-

Na

PAGE 1-B

-(CH<sub>2</sub>)<sub>16</sub>-Me

L7 ANSWER 4 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1994:558704 CAPLUS

DOCUMENT NUMBER: 121:158704

TITLE: A criterion for microphase separation in segmented

polyurethane and polyurethane ureas

AUTHOR(S): Vilensky, V. A.; Lipatov, Y. S.

CORPORATE SOURCE: Inst. Macromol. Chem., Acad. Sci. Ukraine, Kiev,

253160, Ukraine

SOURCE: Polymer (1994), 35(14), 3069-74

CODEN: POLMAG; ISSN: 0032-3861

DOCUMENT TYPE: Journal LANGUAGE: English

AB A criterion for microphase separation in segmented polyurethanes and poly(urethane ureas) was proposed. The existence of correlation between the ratios  $\chi hs/(\chi hs)$  cr and the degree of segregation ( $\alpha seg$ ) was established, where  $\chi hs$  was the thermodn. interaction parameter between soft and hard blocks, calculated from the solubility parameters, and ( $\chi hs$ )cr was its critical value, calculated using the Scott equation. Correlation between the ratio  $\chi hs/(\chi hs)$ cr, the degree of segregation  $\alpha seg$ , and the flexibility parameter  $\sigma$  was also found.

IT 82822-98-8D, derivs., polymers with MDI and polytetramethylene glycol, block 157497-56-8D, (R)-Sulfosuccinic acid dihydrazide, derivs., polymers with MDI and polytetramethylene glycol, block RL: PRP (Properties)

(microphase separation in, calcn. of)

RN 82822-98-8 CAPLUS

CN Butanedioic acid, sulfo-, 1,4-dihydrazide (9CI) (CA INDEX NAME)

RN 157497-56-8 CAPLUS

CN Butanedioic acid, sulfo-, 1,4-dihydrazide, (R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

$$H_{2N}$$
 $H_{2N}$ 
 $H$ 

L7 ANSWER 5 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1994:247810 CAPLUS

DOCUMENT NUMBER: 120:247810

TITLE: Nitrogen Analogs of AOT. Synthesis and Properties AUTHOR(S): Leydet, A.; Boyer, B.; Lamaty, G.; Roque, J. P.;

Catlin, K.; Menger, F. M.

CORPORATE SOURCE: Laboratoire de Chimie Organique Physique, Universite

de Montpellier II, Montpellier, 34095, Fr.

SOURCE: Langmuir (1994), 10(4), 1000-2

CODEN: LANGD5; ISSN: 0743-7463

DOCUMENT TYPE:

LANGUAGE:

Journal English

AB The synthesis of AOT (bis(2-ethylhexyl)sodium sulfosuccinate) analogs, in which the two esters are replaced by more chemical stable amides, is described. The nitrogen analogs of AOT form reverse micelles in chloroform with omax values similar to that of AOT. The compds. are, however, too insol. to form reverse micelles in heptane. Various alkyl groups can be placed on the amide groups of the AOT analogs in order to modulate the hydrophilic/lipophilic balance.

IT 154521-67-2P 154521-68-3P 154521-69-4P

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (preparation and characterization of)

RN 154521-67-2 CAPLUS

CN 2-Butanesulfonic acid, 1,4-bis[(2-ethylhexyl)amino]-1,4-dioxo-, monosodium salt (9CI) (CA INDEX NAME)

## Na

RN 154521-68-3 CAPLUS

CN 2-Butanesulfonic acid, 1,4-bis[ethyl(2-ethylhexyl)amino]-1,4-dioxo-, sodium salt (9CI) (CA INDEX NAME)

### Na

RN 154521-69-4 CAPLUS

CN 2-Butanesulfonic acid, 1,4-bis[(2-ethylhexyl)octylamino]-1,4-dioxo-,

### Na

L7 ANSWER 6 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1992:23082 CAPLUS

DOCUMENT NUMBER: 116:23082

TITLE: Recording ink containing sulfonate dispersant for ink

jet recording

INVENTOR(S): Takimoto, Hiroshi; Kajikawa, Akira; Yoneyama, Tomio

PATENT ASSIGNEE(S): Mitsubishi Electric Corp., Japan

SOURCE: Eur. Pat. Appl., 22 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE	
EP 448055	A2	19910925	EP 1991-104257		19910319	<
EP 448055	<b>A</b> 3	19920805				
EP 448055	B1	19960103				-
R: DE, FR, GB						
JP 03273067	A2	19911204	JP 1990-70089		19900320	<
JP 2952944	В2	19990927				
JP 03287676	A2	19911218	JP 1990-88593		19900403	<
JP 2841678	B2	19981224				
JP 04007372	A2	19920110	JP 1990-108294		19900424	<
JP 2969775	B2	19991102				
JP 04018469	A2	19920122	JP 1990-122501		19900511	<
JP 04039366	A2	19920210	JP 1990-147704		19900606	
JP 2870991	B2	19990317				
US 5125968	A	19920630	US 1991-672554		19910320	<
JP 04213374	A2	19920804	JP 1991-59953		19910325	
JP 2970015	B2	19991102	01 1331 03300		13310020	`
PRIORITY APPLN. INFO.:			JP 1990-70089	A	19900320	
			JP 1990-88593	A		
			JP 1990-108293	A		
			JP 1990-108294	A	19900424	
			JP 1990-122501	A		
				A		
OWNED COUNCE/C).	MADDAM	116.22002	JP 1990-147704	А	19900606	

OTHER SOURCE(S): MARPAT 116:23082

AB Light— and water—resistant inks with good storage stability comprise an aqueous medium, pigment, and ≥1 sulfonate dispersant such as R1CH2CH(OH)(CH2)mSO3M, R2CH:CH(CH2)nSO3M, R3O2CCH(SO3M)CH2CO2R4, MO2CCH(SO3M)CH2CONHR5, R6NHCOCH(SO3M)CH2CONHR6, MO2CCH(SO3M)CH2CO2R7, R8CON(R9)R10SO3M, or R11-p-C6H4O(C2H4O)pSO3M (R1-2 = C8-20 alkyl; R3, R4, R6 = C6-16 alkyl or alkenyl; R5, R7, R8 = C10-20 alkyl or alkenyl; R9 = C1-4 alkyl; R10 = C1-3 alkylene; R11 = R6-18 alkyl; m = 1-3; n = 1-3; p = 1-15; M = Na, NH4). An ink contained PEG 200 15, C.I. Pigment Red 122 4, C12H25CH2CH(OH)CH2SO3Na 1.5, and water 79.5%.

IT 138101-80-1

RL: USES (Uses)

(dispersing agent, for jet printing inks) RN 138101-80-1 CAPLUS CN 2-Butanesulfonic acid, 1,4-bis(octylamino)-1,4-dioxo-, monosodium salt (9CI) (CA INDEX NAME)

### Na

ANSWER 7 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1990:615153 CAPLUS

DOCUMENT NUMBER:

113:215153

TITLE:

Removal of asphalt or resin from hydrocarbons using

both organic solvents and water

INVENTOR(S):

Muller, Alain

PATENT ASSIGNEE(S):

Societe Nationale Elf Aquitaine (SNEA), Fr.

SOURCE:

PCT Int. Appl., 19 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

French

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

	PAT	CENT 1	мо.			KIN	D	DATE	AP	PLICATION NO.		DATE	
	WO	9006: W:	350 JP,	us		A1	<del>-</del>	19900614	WO	1989-FR601		19891123	<
			BE,		GB.	IT,	NL						
	FR	2639	649	•	•	A1		19900601	FR	1988-15387		19881125	<
	FR	2639	649			B1		19910125					
	ΕP	4209	46			<b>A</b> 1		19910410	EP	1989-913186		19891123	<
		R:	BE,	DE,	GB,	IT,	NL						
	JP	03502	2342			Т2		19910530	JP	1990-500209		19891123	<
	CA	2003	833			AA		19900525	CA	1989-2003833		19891124	<
PRIO	RITY	( APP	LN.	INFO	.:				FR	1988-15387	A ·	19881125	
									WO	1989-FR601	W	19891123	

AB Asphalt and/or resin is removed from a hydrocarbon feedstock, e.g., asphalt-containing crude oil, distillation residues, or deasphalted petroleum by solvent extraction using water containing a surfactant, e.g., a sulfonate, and metal salts to sep. the hydrocarbon-solvent emulsion. The mixture seps. into an upper layer of treated hydrocarbons in solvent, a middle layer of water, and a bottom layer containing the asphalt and/or resin. Prior to separation the mixture is agitated for 30 s to 10 min at ambient temperature to 170°.

#### IT 116453-32-8

RL: USES (Uses)

(surfactant, in removal of asphalts and resins from hydrocarbons using organic solvents in water)

116453-32-8 CAPLUS RN

2-Butanesulfonic acid, 1,4-diamino-1,4-dioxo- (9CI) (CA INDEX NAME) CN

L7 ANSWER 8 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1990:614146 CAPLUS

DOCUMENT NUMBER:

113:214146

TITLE: Removal of fillers from wastepaper by flotation in the

presence of sulfonates as surfactants

INVENTOR(S): Behler, Ansgar; Hoefer, Rainer; Hornfeck, Klaus; Von

Rybinski, Wolfgang

Henkel K.-G.a.A., Germany PATENT ASSIGNEE(S):

SOURCE:

Ger. Offen., 5 pp. CODEN: GWXXBX

DOCUMENT TYPE:

Patent

1

LANGUAGE: FAMILY ACC. NUM. COUNT: German

PATENT INFORMATION:

	PAT	TENT NO.			KIN	D DATE	APPLICATION NO.		DATE	
	DE	3900940			A1	19900719	DE 1989-3900940		19890114	<
	WO						WO 1990-EP22		19900105	<
				•	•	NO, US				
							GB, IT, LU, NL, SE			
	AU	9048085			<b>A</b> 1	19900813	AU 1990-48085		19900105	<
						19921029				
							EP 1990-900179		19900105	<
	EP	453449			В1	19930602				
		R: AT	, BE,	CH,	DE,	ES, FR, GB,	IT, LI, NL, SE			
	JP	04502789	9		T2	19920521	JP 1990-501043		19900105	<
	AT	90122			E	19930615	AT 1990-900179		19900105	<
	ES	2041172			Т3	19931101	ES 1990-900179		19900105	
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	CA	2007736			AA	19900714	CA 1990-2007736		19900115	<
	NO	9102125			Α	19910603			19910603	<
	FI	95606			В	19951115			19910709	
		95606				19960226				-
	US	5308448			Α	19940503			19930208	<
P	RIORITY	APPLN.	INFO	. :			DE 1989-3900940		19890114	
							EP 1990-900179			
							WO 1990-EP22		19900105	
							US 1991-721515			
							00 TOOT 15TOTO	דמ	エンフエリノエム	

OTHER SOURCE(S): MARPAT 113:214146

In the title process, filler removal is increased by flotation in the presence of the sulfonates RCH(SO3M1)CO2M2 (R = C6-20 alkyl; M1 = H, alkali metal, NH4; M2 = H, alkali metal, NH4, C1-4 alkyl) or alkali metal or amine salts of sulfonated C12-22 fatty acids, sulfosuccinic acid, or its esters or amides, and/or sec-alkanesulfonic acids. A suspension of 23 g kaolin in 9 L H2O at pH 8.5-9.0 was subjected to flotation in the presence of 0.2 g Na mono-C12-18 alkyl sulfosuccinate, resulting in a 94% removal of kaolin.

116453-32-8D, alkali metal and amine salts

RL: USES (Uses)

(flotation agents, for filler removal from wastepaper)

RN 116453-32-8 CAPLUS

CN 2-Butanesulfonic acid, 1,4-diamino-1,4-dioxo- (9CI) (CA INDEX NAME)

ANSWER 9 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1989:61540 CAPLUS

DOCUMENT NUMBER:

110:61540

TITLE:

Alkyl and alkenyl aspartic acids or their salts in

collectors for flotation of nonsulfide ores

INVENTOR(S):

Kottwitz, Beatrix; Von Rybinski, Wolfgang; Koester,

Rita

PATENT ASSIGNEE(S):

Henkel K.-G.a.A., Fed. Rep. Ger.

SOURCE:

Ger. Offen., 6 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
DE 3641579	A1	19880616	DE 1986-3641579	- <b>-</b> .	19861205 <
EP 270018	A2	19880608	EP 1987-117541		19871127 <
EP 270018	A3	19900418			
EP 270018	B1	19920617			
R: AT, DE, ES,	, FR, GE	, SE			
AT 77262	E	19920715	AT 1987-117541		19871127 <
ES 2031869	Т3	19930101	ES 1987-117541		19871127 <
FI 8705336	Α	19880606	FI 1987-5336		19871203 <
FI 84321	В	19910815			
FI 84321	С	19911125			
CN 87107280	Α	19880615	CN 1987-107280		19871203 <
CN 1011296	В	19910123			
US 4790932	Α	19881213	US 1987-128303		19871203 <
AU 8782109	A1	19880609	AU 1987-82109		19871204 <
AU 601244	B2	19900906			
BR 8706570	Α	19880712	BR 1987-6570		19871204 <
ZA 8709141	Α	19880727	ZA 1987-9141		19871204 <
CA 1320769	A1	19930727	CA 1987-553595		19871204 <
PRIORITY APPLN. INFO.:			DE 1986-3641579	Α	19861205
			EP 1987-117541	Α	19871127

AB Flotation with collectors containing N-alkyl aspartic and N-alkenyl aspartic acids and their salts is suitable for higher yields at equal amts. and selectivity, or equal yields at lower collector concns. Thus, in flotation of scheelite ore the collector consisted of 2:1 weight mixture of tallow ammine-derived sulfosuccinamide and the Na salts of N-C16-18-alkylaspartic acid used at 500 g/ton ore. The resulting concentrate contained WO3 28.3, CaO 15.8, SiO2 21.1, vs. 10.6, 8.6, and 34.8% resp. for a conventional collector.

IT 116453-32-8D, tallow-alkyl derivs.

RL: PROC (Process)

(flotation collectors, anionic, with alkyl- and alkenylaspartic acid and salts, for nonsulfide ores)

RN 116453-32-8 CAPLUS

CN 2-Butanesulfonic acid, 1,4-diamino-1,4-dioxo- (9CI) (CA INDEX NAME)

L7 ANSWER 10 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1988:532575 CAPLUS

DOCUMENT NUMBER: 109:132575

TITLE: Surfactant mixtures as collectors in flotation of

nonsulfidic ores

INVENTOR(S): Koester, Rita; Von Rybinski, Wolfgang

PATENT ASSIGNEE(S): Henkel K.-G.a.A., Fed. Rep. Ger.

SOURCE: Ger. Offen., 8 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 3641447	A1	19880609	DE 1986-3641447	19861204 <

	EP 270933	A2	19880615	EP 1987-117456	19871126 <				
	EP 270933	A3	19891025						
	EP 270933	В1	19920722						
•		, ES, FR, GE							
	US 4790931	Α	19881213	US 1987-127749	19871202 <				
	FI 8705335	Α	19880605	FI 1987-5335	19871203 <				
	FI 83044	В	19910215						
	FI 83044	С	19910527						
	AU 8782066	A1	19880609	AU 1987-82066	19871203 <				
	AU 598069	B2	19900614						
	CN 87107281	Α	19880615	CN 1987-107281	19871203 <				
	CN 1012420	В	19910424						
	ZA 8709095	Α	19880727	ZA 1987-9095	19871203 <				
	BR 8706550	A	19880712	BR 1987-6550	19871204 <				
PRIC	RITY APPLN. INFO			DE 1986-3641447	A 19861204				
AB				alc. polyglycol eth					
				n flotation of nons					
	Thus, scheelite ore powder (containing WO3 0.3, CaO 8.8, and SiO2 55.8%)								
	having particle size <200 $\mu m$ was processed using a 2:1 mixture of an								
	anionic and a nonionic surfactants. The anionic component was Na salt of								
				vamine, and the no					
				d on C12-18 fatty a					
				was water glass a					
				ditioning for 10 mi					
				.9.5. Conditionin					
					13.3, CaO 32.9, and SiO2				
				for a conventional	collector at				
4	.apprx.40% high			,					
IT				692-36-5D, Sodium					
	sulfosuccinamid		mine-derived	<b>.</b>					
	RL: PROC (Proce	•							
				on collectors with	end				
	group-termin		arc. boradra	col ethers)					
RN	116453-32-8 CA								
CN	2-Butanesulfoni	.c acid, 1,4	-diamino-1,4	l-dioxo- (9CI) (CA	. INDEX NAME)				
	O SO3H O								

RN 116692-36-5 CAPLUS

CN 2-Butanesulfonic acid, 1,4-diamino-1,4-dioxo-, monosodium salt (9CI) (CA INDEX NAME)

Na

L7 ANSWER 11 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1985:160051 CAPLUS

DOCUMENT NUMBER:

102:160051

TITLE:

Preparation of surfactants with demonstrated

pharmacological activity

AUTHOR(S):

Kabachnyi, V. I.; Chernykh, V. P.; Kabachnyi, G. I.;

Sopel'nik, E. M.

CORPORATE SOURCE:

Khar'k. Farm. Inst., Kharkov, USSR

SOURCE:

Khimiko-Farmatsevticheskii Zhurnal (1985),

19(1), 43-6

CODEN: KHFZAN; ISSN: UU23-1134

DOCUMENT TYPE:

LANGUAGE:

Journal Russian

OTHER SOURCE(S):

CASREACT 102:160051

AB Sixteen surfactant sulfosuccinic acid heterylamides were prepared and tested for pharmacol. activity and toxicity in mice. Several of the compds. exhibited anti-inflammatory activity comparable to that of butadione, and several caused lowering of blood sugar levels comparable to those produced by butamide.

IT 95896-38-1 95896-39-2 95896-40-5 95896-41-6 95896-42-7 95896-43-8

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(pharmacol. of)

RN 95896-38-1 CAPLUS

CN 2-Butanesulfonic acid, 1-amino-1,4-dioxo-4-(1,3,4-thiadiazol-2-ylamino)-(9CI) (CA INDEX NAME)

RN 95896-39-2 CAPLUS

CN 2-Butanesulfonic acid, 1-amino-1,4-dioxo-4-[(5-propyl-1,3,4-thiadiazol-2-yl)amino]- (9CI) (CA INDEX NAME)

RN 95896-40-5 CAPLUS

CN 2-Butanesulfonic acid, 1-amino-4-[(5-butyl-1,3,4-thiadiazol-2-yl)amino]-1,4-dioxo-(9CI) (CA INDEX NAME)

RN 95896-41-6 CAPLUS

CN 2-Butanesulfonic acid, 1-(methylamino)-1,4-dioxo-4-[(5-propyl-1,3,4-thiadiazol-2-yl)amino]- (9CI) (CA INDEX NAME)

2-Butanesultonic acid, 1-(methylamino)-4-[[5-(1-methylethyl)-1,3,4-thiadiazol-2-yl]amino]-1,4-dioxo-(9CI) (CA INDEX NAME)

RN 95896-43-8 CAPLUS

CN

CN 2-Butanesulfonic acid, 1-(methylamino)-4-[[5-(2-methylpropyl)-1,3,4-thiadiazol-2-yl]amino]-1,4-dioxo-(9CI) (CA INDEX NAME)

IT 95896-22-3P 95896-23-4P 95896-24-5P 95896-25-6P 95896-26-7P 95896-27-8P

RL: BAC (Biological activity or effector, except adverse); BPR (Biological process); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); PROC (Process); USES (Uses)

(preparation and pharmacol. of)

RN 95896-22-3 CAPLUS

CN 2-Butanesulfonic acid, 1-amino-1,4-dioxo-4-(1,3,4-thiadiazol-2-ylamino)-, monosodium salt (9CI) (CA INDEX NAME)

Na

RN 95896-23-4 CAPLUS

CN 2-Butanesulfonic acid, 1-amino-1,4-dioxo-4-[(5-propyl-1,3,4-thiadiazol-2-yl)amino]-, monosodium salt (9CI) (CA INDEX NAME)

Na

2-Butanesultonic acid, 1-amino-4-[(5-butyl-1,3,4-thiadiazol-2-yl)amino]-1,4-dioxo-, monosodium salt (9CI) (CA INDEX NAME)

Na

RN 95896-25-6 CAPLUS

CN

CN 2-Butanesulfonic acid, 1-(methylamino)-1,4-dioxo-4-[(5-propyl-1,3,4-thiadiazol-2-yl)amino]-, monosodium salt (9CI) (CA INDEX NAME)

Na

RN 95896-26-7 CAPLUS

CN 2-Butanesulfonic acid, 1-(methylamino)-4-[[5-(1-methylethyl)-1,3,4-thiadiazol-2-yl]amino]-1,4-dioxo-, monosodium salt (9CI) (CA INDEX NAME)

Na

RN 95896-27-8 CAPLUS

CN 2-Butanesulfonic acid, 1-(methylamino)-4-[[5-(2-methylpropyl)-1,3,4-thiadiazol-2-yl]amino]-1,4-dioxo-, monosodium salt (9CI) (CA INDEX NAME)

Na

CAPLUS COPYRIGHT 2006 ACS on STN ANSWER 12 OF 19

ACCESSION NUMBER:

1984:439598 CAPLUS

DOCUMENT NUMBER:

101:39598

TITLE:

Synthesis of ionomeric polyurethane latexes

AUTHOR(S):

Sukhorukova, A. S.; Grekov, A. P.; Levchenko, N. I.;

Navrotskaya, R. P.

CORPORATE SOURCE:

Inst. Khim. Vysokomol. Soedin., Kiev, USSR

SOURCE:

Sint. Iskusstv. Lateksy: Poluch. Modif., Mater. Vses.

Lateksnoi Konf., 6th (1982), Meeting Date 1981, 115-20. Editor(s): Tikhomirov, G. S.

TsNIITEneftekhim: Moscow, USSR.

CODEN: 51NMA3

DOCUMENT TYPE:

Conference

LANGUAGE: Russian

AB Ionomeric urethane rubber latexes were prepared by reaction of poly(propylene oxide)glycol or poly(tetramethylene oxide)glycol (I) with tolylene diisocyanate (II), followed by chain extension with alkylmalonic or thioalkylsuccinic acid dihydrazides. The latexes formed transparent, elastic films, whose tensile strength and modulus of elasticity increased with increasing substituted dihydrazide concentration Alternatively, cationic polyurethane latexes were prepared by reaction of I with II to form a prepolymer, which was dissolved in DMF-Me2CO mixture, followed by chain extension with aqueous dihydrazide solns. containing tertiary ammonium groups in the side chain. Anionic polyurethane latexes were prepared by using hydrophobic organic solvents, e.g., PhMe at the chain extension stage. physicomech. properties and uses of the latexes were discussed. IT

77986-50-6D, ionic derivs.

RL: USES (Uses)

(rubber, latexes)

77986-50-6 CAPLUS RN

CN Butanedioic acid, sulfo-, 1,4-dihydrazide, monosodium salt, polymer with 1,3-diisocyanatomethylbenzene and  $\alpha$ -hydro- $\omega$ -hydroxypoly(oxy-1,4-butanediyl) (9CI) (CA INDEX NAME)

CM 1

CRN 66693-73-0 CMF C4 H10 N4 O5 S . Na

Na

CM 2

26471-62-5 CRN CMF C9 H6 N2 O2 CCI IDS

25190-06-1 CRN

(C4 H8 O)n H2 O CMF

CCI PMS

ANSWER 13 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1982:492899 CAPLUS

DOCUMENT NUMBER:

97:92899

TITLE:

Ionic polyacylurethane semicarbazides

AUTHOR(S):

Sukhorukova, S. A.

CORPORATE SOURCE:

Inst. Khim. Vysokomol. Soedin., Kiev, USSR

SOURCE:

Sint. Poliuretanov (1981), 77-82.

Editor(s): Omel'chenko, S. I. Izd. Naukova Dumka:

Kiev, USSR. CODEN: 48BKA9

DOCUMENT TYPE: Conference LANGUAGE: Russian

Ionic polyacylurethane semicarbazide dispersions were prepared by polymerization of AB dicarboxylic acid dihydrazides with polytetramethylene glycol (mol. weight 1000) and tolylene diisocyanate in Me2CO or DMF. The ionic dispersions are stable for 6 mo and readily form elastic films having enhanced hydrophilicity.

IT 82822-99-9P

> RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of ionic dispersions of)

RN82822-99-9 CAPLUS

CN Butanedioic acid, sulfo-, 1,4-dihydrazide, polymer with 1,3-diisocyanatomethylbenzene and  $\alpha$ -hydro- $\omega$ -hydroxypoly(oxy-1,4-butanediyl) (9CI) (CA INDEX NAME)

CM 1

CRN 82822-98-8 CMF C4 H10 N4 O5 S

CM

26471-62-5 CRN CMF C9 H6 N2 O2 CCI IDS

CM

25190-06-1 (C4 H8 O)n H2 O CMF

CCI PMS

CRN

1.7 ANSWER 14 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1981:408393 CAPLUS

DOCUMENT NUMBER: 95:8393

TITLE: Synthesis of anion-active polyurethane ionomers

AUTHOR(S): Sukhorukova, S. A.; Levchenko, N. I.; Grekov, A. P.

CORPORATE SOURCE: Inst. Khim. Vysokomol. Soedin., Kiev, USSR

SOURCE: Ukrainskii Khimicheskii Zhurnal (Russian Edition) (

1981), 47(3), 286-90

CODEN: UKZHAU; ISSN: 0041-6045

DOCUMENT TYPE: Journal LANGUAGE: Russian

Dihydrazides containing SO3Na groups in the side chains are used as chain extenders in the preparation of water-dispersible polyurethane ionomers for finishing fibers, leather, wood, paper, and other materials. The optimum conditions for preparation of the ionomers as aqueous dispersions were examined based on the dependence of properties of the systems and their films on the ionic center concentration, urethane segment length, dihydrazide and solvent nature, and dispersion method. The properties of the polymer dispersions prepared in PhMe depended significantly on the dispersing method. The optimum concentration of anionic groups in the polymer was 6%. The properties of polyurethanes prepared from poly(diethylene glycol adipate) and from polytetramethylene glycol (I) at an optimum content of ionic centers were similar. The most effective solvent for preparation of the ionomers was DMF. Polymers based on 1,6-hexamethylene diisocyanate and I had better mech. properties than TDI-based polymers.

IT 77866-24-1P 77866-26-3P 77884-42-5P

77974-01-7P 77986-50-6P

RL: SPN (Synthetic preparation); PREP (Preparation)

(ionomer, preparation and properties of)

RN 77866-24-1 CAPLUS

Butanedioic acid, sulfo-, 1,4-dihydrazide, monosodium salt, polymer with CN 1,6-diisocyanatohexane and  $\alpha$ -hydro- $\omega$ -hydroxypoly(oxy-1,4-

butanediyl) (9CI) (CA INDEX NAME)

CM

CRN 66693-73-0

CMF C4 H10 N4 O5 S . Na

SO3H H2N-NH-C-CH-CH2-C-NH-NH2

Na

CM 2

CRN 25190-06-1 CMF (C4 H8 O)n H2 O CCI PMS

HO (CH<sub>2</sub>) 
$$_4$$
 - O  $_n$ 

CM 3

CRN 822-06-0 CMF C8 H12 N2 O2

OCN- (CH2)6-NCO

RN 77866-26-3 CAPLUS

CN Hexanedioic acid, dihydrazide, polymer with 1,6-diisocyanatohexane,  $\alpha$ -hydro- $\omega$ -hydroxypoly(oxy-1,4-butanediyl) and sulfobutanedioic acid 1,4-dihydrazide, monosodium salt (9CI) (CA INDEX NAME)

CM 1

CRN 66693-73-0 CMF C4 H10 N4 O5 S . Na

Na

CM 2

CRN 25190-06-1 CMF (C4 H8 O)n H2 O CCI PMS

HO 
$$\left[ (CH_2)_4 - O \right]_n$$

CM 3

CRN 1071-93-8 CMF C6 H14 N4 O2

CM 4

CRN 822-06-0 CMF C8 H12 N2 O2

OCN-(CH<sub>2</sub>)<sub>6</sub>-NCO

RN 77884-42-5 CAPLUS

CN 1,3-Benzenedicarboxylic acid, dihydrazide, polymer with
1,6-diisocyanatohexane, α-hydro-ω-hydroxypoly(oxy-1,4butanediyl) and sulfobutanedioic acid 1,4-dihydrazide monosodium salt
(9CI) (CA INDEX NAME)

CM 1

CRN 66693-73-0 CMF C4 H10 N4 O5 S . Na

Na

CM 2

CRN 25190-06-1 CMF (C4 H8 O)n H2 O CCI PMS

HO 
$$\left[ (CH_2)_4 - O \right]_n$$

CM 3

CRN 2760-98-7 CMF C8 H10 N4 O2

$$\begin{array}{c|c} & & & \\ \mathbf{H_2N-NH-C} & & & \mathbf{C-NH-NH_2} \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ \end{array}$$

CM 4

CRN 822-06-0 CMF C8 H12 N2 O2

OCN-(CH<sub>2</sub>)<sub>6</sub>-NCO

RN 77974-01-7 CAPLUS

CN Hexanedioic acid, polymer with 1,3-diisocyanatomethylbenzene, 2,2'-oxybis[ethanol] and sulfobutanedioic acid 1,4-dihydrazide monosodium salt (9CI) (CA INDEX NAME) CM 1 CRN 66693-73-0 CMF C4 H10 N4 O5 S . Na SO3H H2N-NH-C-CH-CH2-C-NH-NH2 Na CM 2 CRN 26471-62-5 CMF C9 H6 N2 O2 CCI IDS NCO OCN D1-Me CM 3 CRN 124-04-9 CMF C6 H10 O4  $HO_2C-(CH_2)_4-CO_2H$ CM 4 CRN 111-46-6 CMF C4 H10 O3 но-сн2-сн2-о-сн2-сн2-он RN 77986-50-6 CAPLUS

RN 77986-50-6 CAPLUS
CN Butanedioic acid, sulfo-, 1,4-dihydrazide, monosodium salt, polymer with 1,3-diisocyanatomethylbenzene and α-hydro-ω-hydroxypoly(oxy-1,4-butanediyl) (9CI) (CA INDEX NAME)

CM 1

CRN 66693-73-0 CMF C4 H10 N4 O5 S . Na

Na

CM 2

CRN 26471-62-5 CMF C9 H6 N2 O2

CCI IDS

D1-Me

CM 3

CRN 25190-06-1

CMF (C4 H8 O)n H2 O

CCI PMS

HO (CH<sub>2</sub>) 
$$_4$$
 - O  $_n$ 

L7 ANSWER 15 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1981:55848 CAPLUS

DOCUMENT NUMBER:

94:55848

TITLE:

Direct positive image formation process

PATENT ASSIGNEE(S):

Konishiroku Photo Industry Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 13 pp. CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

. 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 55062443	A2	19800510	JP 1978-136207	19781102 <
JP 58028570	В4	19830616		

PRIORITY APPLN. INFO.:

JP 1978-136207 A 19781102

Direct-pos. type photog. materials having internal latent image type emulsion layers are imagewise exposed and developed in the presence of a fogging agent selected from RNHNHZZ1kZ2lNHNHR1, R2NHNHZ3Z4mCR3:NNHR4, and R5NHN:CR6Z5nCR7:NNHR8 (R, R1, R2, R4, R5, R8 = aryl, heterocyclic moiety; Z, Z2, Z3 = CO, SO2; R3, R6, R7 = H, lower alkyl, aryl; Z1, Z4, Z5 = divalent organic moiety; R3 and R7 may combine with Z4 and Z5, resp., to form 5- or 6-membered rings; k, l, m, n = 0, 1). Thus, a fogging agent p-HO3SC6H4NHNHCOCONHNHC6H4SO3H-p mg/mol Ag halide was added to an internal

latent image type Ag(Br,Cl,I) emulsion and the emulsion was coated on a film support. The photog. film was then imagewise exposed and developed to give Dmax and Dmin of 0.85 and 0.11, resp., vs. 0.08 and 0.07, resp., for a fogging agent-free control.

IT 70794-87-5

RL: USES (Uses)

(photog. fogging agent, for direct-pos. emulsions)

RN 70794-87-5 CAPLUS

CN Butanedioic acid, sulfo-, 1,4-bis(2-phenylhydrazide) (9CI) (CA INDEX NAME)

L7 ANSWER 16 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1979:466218 CAPLUS

DOCUMENT NUMBER: 91:66218

TITLE: Direct-positive photographic products

AUTHOR(S): Anon. CORPORATE SOURCE: UK

SOURCE: Research Disclosure (1979), 181, 246 (No.

18171)

CODEN: RSDSBB; ISSN: 0374-4353

DOCUMENT TYPE: Journal; Patent

LANGUAGE: English

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

RD 181071 19790510

PRIORITY APPLN. INFO.:

RD 1979-181071 19790510

AB Hydrazide derivs. are described which can efficiently function as fogging agents with smaller quantities and at lower pH values than those known. These compds., which are especially useful in direct-pos. photog. products, have the formulas R1NHNHZ(Z1)k(Z2)l=NHNHR2, R3NHNHZ4(Z5)mCR4=NNHR5, and R6NHNHCR7=(Z6)n=CR8=NNHR9 (R1,R2,R3,R5,R6,R9 = aryl or heterocycle; R4,R7,R8 = H, alkyl, or aryl; Z,Z2,Z4 = CO, SO2; Z1,Z5,Z6 = a divalent organic group; k,l,m, or n = 0 or 1).

IT 70794-87-5

SOURCE:

RL: USES (Uses)

(fogging agent, for direct-pos. photog. materials)

RN 70794-87-5 CAPLUS

CN Butanedioic acid, sulfo-, 1,4-bis(2-phenylhydrazide) (9CI) (CA INDEX NAME)

L7 ANSWER 17 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1978:425321 CAPLUS

DOCUMENT NUMBER: 89:25321

TITLE: Hydrazides of sulfodicarboxylic acid sodium salts as

monomers for aqueous dispersion of polyurethanes

INVENTOR(S): Sukhorukova, S. A.; Levchenko, N. I.; Klimenko, N. S.;

Grekov, A. P.

PATENT ASSIGNEE(S): Institute of the Chemistry of High-Molecular-Weight

Compounds, Academy of Sciences, Ukrainian S.S.R, USSR U.S.S.R. From: Otkrytiya, Izobret., Prom. Obraztsy,

Tovarnye Znaki 1978, 55(10), 77.

CODEN: URXXAF

DOCUMENT TYPE: LANGUAGE:

Patent Russian

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PRIORITY APPLN. INFO.: SU 1976-2418557 A 19761109

AB H2NNHCOXCONHNH2 [(X = CH2CH2CH2CH2SO3Na) [66693-73-0] or

CH(CH2CH2CH2SO3Na) [66693-74-1]] are monomers for aqueous dispersion of

polyurethanes. IT 66693-73-0

IT 66693-73-0 RL: USES (Uses)

(monomers, for aqueous dispersion of polyurethanes)

RN 66693-73-0 CAPLUS

CN Butanedioic acid, sulfo-, 1,4-dihydrazide, monosodium salt (9CI) (CA

INDEX NAME)

### Na

L7 ANSWER 18 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1969:503016 CAPLUS

DOCUMENT NUMBER:

71:103016

TITLE:

SOURCE:

Sizing of polyamide warps of polyfilament yarns

AUTHOR(S): Vinea, E.; Radulescu, Cecilia

CORPORATE SOURCE:

Tesatoria Relon "Panduri", Bucharest, Rom.

Industria Textila (Bucharest, 1950-1973) (1969

), 20(6), 400-2

CODEN: INTBA7; ISSN: 0019-7750

DOCUMENT TYPE: LANGUAGE:

Journal Romanian

AB Sizing expts. with Vinarol DT, THM Schkopau 45/02, Sizing TD, Sopronyl PAA 10-40, and Sopromine 1686 in varying concns. with and without glycerol were conducted to obtain the best recipe for sizing Relon warps. Best results were obtained with a recipe comprising 2.5% Sopronyl PAA 10-40 and 0.3% Sopromine 1686. Comparison of recipes comprising 1.5% Aracet APV [poly-(vinyl alc.)] and 0.5% glycerol or 2% Aracet APV and 0.8% glycerol with recipes containing Vinarol DT showed that the Romanian products were satisfactory but formed more rigid films.

IT 94200-33-6, Sopromine 1686

RL: USES (Uses)

(in sizing of nylon warps)

RN 94200-33-6 CAPLUS

CN Octadecanoic acid, (1,4-dioxo-2-sulfo-1,4-butanediyl)bis(imino-2,1-ethanediyl) ester, monosodium salt (9CI) (CA INDEX NAME)

# - (CH<sub>2</sub>)<sub>16</sub>- Me

L7 ANSWER 19 OF 19 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1951:59473 CAPLUS

DOCUMENT NUMBER: 45:59473
ORIGINAL REFERENCE NO.: 45:10111b-c

TITLE: Direct positive photographs from hydrazine-containing

developers

INVENTOR(S): Ives, Charles E. PATENT ASSIGNEE(S): Eastman Kodak Co.

DOCUMENT TYPE: Patent LANGUAGE: Unavailable

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE					
	US 2563785		19510807	US 1950-159150	19500429 <					
AB	Direct positive ima	ges are	obtained by	exposing an internal	latent-image					
	emulsion to actinic	light,	then develo	ping in a Ag halide de	veloping solution					
	containing a N2H4 compound of the general formula R2NNR2, in which at least 2									
	R's but less than 4 are H and the remaining R's are aryl, aralkyl, acyl,									
	or carboxylic acid amide groups. Suitable compds. are:									
				ulfonamido)ethyl]pheny	lhydrazine,					
				Na sulfosuccinic acid						
IT				ydrazide monosodium sa						
	(in photography)	-	,	-						
RN	66693-73-0 CAPLUS									
CN	Butanedioic acid, s	ulfo-,	1,4-dihvdraz	ide, monosodium salt (	9CI) (CA					
	INDEX NAME)	·	•	•	, , , , , , , , , , , , , , , , , , , ,					

Na

Structure attributes must be viewed using STN Express query preparation.

=> s 18 full

### REG1stRY INITIATED

Substance data SEARCH and crossover from CAS REGISTRY in progress... Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

204 ANSWERS

FULL SEARCH INITIATED 15:21:50 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 1182 TO ITERATE

100.0% PROCESSED 1182 ITERATIONS

SEARCH TIME: 00.00.01

L9 204 SEA SSS FUL L8

L10 295 L9

 $\Rightarrow$  s 110 and py<2002

21808282 PY<2002

L11 224 L10 AND PY<2002

=> s 111 and composition

649375 COMPOSITION

L12 10 L11 AND COMPOSITION

=> s l11 and (quaternary ammonium or quaternary phoshonium)

124926 QUATERNARY

361380 AMMONIUM

61727 QUATERNARY AMMONIUM

(QUATERNARY (W) AMMONIUM)

124926 QUATERNARY

20 PHOSHONIUM

1 QUATERNARY PHOSHONIUM

(QUATERNARY (W) PHOSHONIUM)

L13 5 L11 AND (QUATERNARY AMMONIUM OR QUATERNARY PHOSHONIUM)

=> d 1-5 ibib abs hitstr

L13 ANSWER 1 OF 5 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1999:182730 CAPLUS

DOCUMENT NUMBER:

130:274069

TITLE:

Charge-controlling agent, and electrostatographic developer toner, powder coating for electrostatic

coating, and charging material using it

INVENTOR(S):

Tsuruhara, Toru; Sugata, Kazuaki

PATENT ASSIGNEE(S):

Orient Chemical Industries, Ltd., Japan

SOURCE:

Jpn. Kokai Tokkyo Koho, 22 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent Japanese

LANGUAGE:

. Japa

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
				<del>-</del>
JP 11072969	A2	19990316	JP 1997-249606	19970829 <
OPTTY APPIN INFO .			JP 1997-249606	19970829

OTHER SOURCE(S):

MARPAT 130:274069

AB The charge-controlling agent comprises ≥1 salts kA+.B-k (B-k = benzenesulfonic acid derivative anion or naphthalenesulfonic acid derivative anion). The electrostatog. toner, the powder coating, and the charging material using the agent are also claimed. The agent shows good dispersibility in polymers and high thermal stability.

IT 221388-45-0 221388-75-6 221388-84-7

RL: TEM (Technical or engineered material use); USES (Uses) (charge-controlling agent for electrostatog. developer toner and powder coating)

RN 221388-45-0 CAPLUS

CN Benzenemethanaminium, N,N-dibutyl-N-1-propenyl-, salt with 3-sulfo-1,2-benzenedicarboxylic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 221388-44-9 CMF C8 H5 O7 S

CM 2

CRN 221388-43-8 CMF C18 H30 N

RN 221388-75-6 CAPLUS

CN 1-Butanaminium, N,N,N-tributyl-, salt with 3-sulfo-1,2-benzenedicarboxylic acid (1:1) (9CI) (CA INDEX NAME)

CM :

CRN 221388-44-9

CM 2

CRN 10549-76-5 CMF C16 H36 N

RN 221388-84-7 CAPLUS

CN Benzenemethanaminium, N,N-dibutyl-N-(4-fluorobutyl)-4-(trifluoromethyl)-, salt with 5-sulfo-1,2,4-benzenetricarboxylic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 221388-83-6 CMF C9 H5 O9 S

CM 2

CRN 221388-82-5 CMF C20 H32 F4 N

$$CH_2 - N^{+}$$
 (CH<sub>2</sub>)<sub>4</sub> - F
 $n-Bu$ 
 $n-Bu$ 

L13 ANSWER 2 OF 5 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1990:593403 CAPLUS

DOCUMENT NUMBER:

113:193403

TITLE:

Finishing of fabrics by cationic or amphoteric agents

INVENTOR(S):

Nakao, Katsuaki; Sato, Koji; Ishido, Kazutaka

PATENT ASSIGNEE(S):

Ipposha Oil and Industries Co., Ltd., Japan

SOURCE:

Jpn. Kokai Tokkyo Koho, 5 pp.

SOURCE.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

Japanese LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE \_\_\_\_ \_\_\_\_\_ -----JP 02080665 A2 19900320 JP 1988-233312 19880918 <--

PRIORITY APPLN. INFO.:

JP 1988-233312 19880918

Fabrics are anionized and finished with cationic or amphoteric agents for good durability as a result of chemical reaction. Thus, a cotton fabric was impregnated with a 20% aqueous solution of 1:1 NaHSO3-epichlorohydrin adduct and NaOH at room temperature for 1 min, squeezed, dried at 110° for 10 min, neutralized, washed, dried, impregnated with an aqueous solution of 5 g/L dimethyldistearylammonium chloride at 40° for 30 min, squeezed, and dried at 100° for 10 min to give a fabric with good retention of softness after repeated washing.

130231-16-2 IT

RL: USES (Uses)

(anionizing agents, for fabrics for cationic or amphoteric finishing)

RN 130231-16-2 CAPLUS

CN 1,2-Benzenedicarboxylic acid, 4-sulfo-, polymer with 1,2-ethanediol (9CI) (CA INDEX NAME)

CM 1

CRN 107-21-1 CMF C2 H6 O2

 $HO-CH_2-CH_2-OH$ 

CM 2

CRN 89-08-7 CMF C8 H6 O7 S

L13 ANSWER 3 OF 5 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1978:430591 CAPLUS

DOCUMENT NUMBER:

89:30591

TITLE:

Copolyester hair conditioners

INVENTOR(S):

Quack, Jochen M.; Reng, Alwin; Engelhardt, Friedrich;

Hintermeier, Karl

PATENT ASSIGNEE(S):

Hoechst A.-G., Fed. Rep. Ger.

SOURCE:

Ger. Offen., 60 pp. CODEN: GWXXBX

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 2633418	<b>A</b> 1	19780126	DE 1976-2633418	19760724 <

DE 2633418	B2	19790125			
NL 7708019	Α	19780126	NL 1977-8019		19770719 <
· US 4150216	Α	19790417	US 1977-817054		19770719 <
SE 7708408	Α	19780125	SE 1977-8408		19770721 <
BR 7704834	Α	19780404	BR 1977-4834		19770722 <
ZA 7704435	Α	19780628	ZA 1977-4435		19770722 <
JP 53015437	A2	19780213	JP 1977-87905		19770723 <
BE 857130	A1	19780125	BE 1977-179617		19770725 <
FR 2358878	A1	19780217	FR 1977-22778		19770725 <
AU 7727230	A1	19790125	AU 1977-27230		19770727 <
PRIORITY APPLN. INFO.:			DE 1976-2633418	Α	19760724

Water-soluble hair conditioners contained branched copolyesters of apparent mol. weight 600-5000 and containing SO3M groups (M = alkali metal, NH4, quaternary ammonium salt). The copolyester residues consisted of -COXCO-,-COX1(CO)n+2-, -OX2O-, -OX3On+2- (X = bond, divalent aliphatic, cycloaliph., aromatic optionally containing SO3M; X1 = aliphatic, cycloaliph., aromatic optionally containing SO3M; X2 = divalent aliphatic, cycloaliph., araliph optionally containing SO3M; X3 = aliphatic, cycloaliph. optionally containing SO3M; n = 0-2). Isophthalic acid 311, di-Me isophthalate 5-Na sulfonate 111, pyromellitic dianhydride 54.5, and diethylene glycol 265 g were heated under N to give a copolyester of apparent mol. weight 700-1000. A hair setting lotion consisted of 3 g copolyester, 46.8 g isopropanol, and 0.2 g perfume.

IT 65408-81-3

AΒ

RL: BIOL (Biological study)
 (for hair conditioners)

RN 65408-81-3 CAPLUS

CN 1,2,4-Benzenetricarboxylic acid, 5-sulfo-, 1,2,4-trimethyl ester, sodium salt, polymer with dimethyl 1,3-benzenedicarboxylate, dimethyl 1,4-benzenedicarboxylate and 2,2'-oxybis[ethanol] (9CI) (CA INDEX NAME)

CM 1

CRN 65408-80-2 CMF C12 H12 O9 S . Na

Na

CM 2

CRN 1459-93-4 CMF C10 H10 O4

CM 3

CRN 120-61-6 CMF C10 H10 O4

CM 4

CRN 111-46-6 CMF C4 H10 O3

но-сн2-сн2-о-сн2-сн2-он

L13 ANSWER 4 OF 5 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1974:425382 CAPLUS

DOCUMENT NUMBER:

81:25382

TITLE:

3,5-Bis ( $\beta$ -hydroxyethoxycarbonyl) benzenesulfonic

acid alkali metal salts

INVENTOR(S):

Terasawa, Isao; Ogura, Sei; Tanaka, Tatsundo;

Nakamura, Itaru

PATENT ASSIGNEE(S):

Toray Industries, Inc.

SOURCE:

Jpn. Kokai Tokkyo Koho, 5 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
JP 49031634	A2	19740322	JP 1972-72560	19720721 <	
JP 50013252	B4	19750519			
PRIORITY APPLN. INFO.:			JP 1972-72560 A	19720721	

Title salts were prepared by esterification of 5-MO3SC6H3(CO2H)2-1,3 (M = alkali metal) with HOCH2CH2OH in the presence of quaternary ammonium or alkali metal compds., e.g., Et4NOH, LiOAc, Na3PO4, or NaO2C(CH2)8CO2Na.

IT 33562-89-9

RL: RCT (Reactant); RACT (Reactant or reagent)
 (esterification with ethylene glycol)

RN 33562-89-9 CAPLUS

CN 1,2-Benzenedicarboxylic acid, 4-sulfo-, monosodium salt (9CI) (CA INDEX NAME)

### Na

L13 ANSWER 5 OF 5 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1966:84517 CAPLUS

DOCUMENT NUMBER: 64:84517 ORIGINAL REFERENCE NO.: 64:15858a-c

TITLE: Alkyl isoquinolinium salts of aromatic carboxylic

acids

INVENTOR(S): Wakeman, Reginald L.; Coates, Joseph F.

PATENT ASSIGNEE(S): Millmaster Onyx Corp.

SOURCE: DOCUMENT TYPE: Patent LANGUAGE: Unavailable

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

DATE PATENT NO. KIND APPLICATION NO. DATE ---------------19660215 US 3235556 US 1963-262836 19630305 <--US PRIORITY APPLN. INFO.: 19630305 The title compds. were prepared by the reaction of N-alkyl isoquinolinium quaternary ammonium compds. having 8-18 C atoms in the alkyl radical with the free acid or salts of aromatic mono-, di-, or polycarboxylic acids. The compds. were shown to be active against Staphylococcus aureus, Salmonella typhosa, and Aspergillus niger, and exhibited low H2O solubility, generally not in excess of 3 parts by weight/100 parts solution at 22°. Thus, from a stock solution containing 10 weight-% sodium benzoate there was taken an aliquot containing 0.035 equivalent of BzONa, a chemical equivalent amount of a com. grade lauryl isoquinolinium bromide in the form of a 10 weight-% solution was added to the agitated solution, the mixture poured into a separatory funnel, and the organic layer dried in vacuo to give in 90% yield lauryl isoquinolinium benzoate. Similarly prepared were the following: octyl isoquinolinium benzoate, di(lauryl isoquinolinium) terephthalate, tetra(lauryl isoquinolinium) pyromellitate, tetra(myristyl isoquinolinium) pyromellitate, tetra(cetyl isoquinolinium) pyromellitate, tetra(stearyl isoquinolinium) pyromellitate, and lauryl isoquinolinium toluate. IT 5201-73-0, Isoquinolinium, 2-dodecyl-, 4-sulfophthalate (3:1)

(as bactericide)

5201-73-0 CAPLUS RN

CN Isoquinolinium, 2-dodecyl-, 4-sulfophthalate (3:1) (8CI) (CA INDEX NAME)

CM 1

CRN 46687-30-3 CMF C8 H3 O7 S

CM 2

CRN 16826-19-0 CMF C21 H32 N

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(Falle 'HOME' ENTERED AT 14:58:23 ON 16 MAR 2006)
     FILE 'CAPLUS' ENTERED AT 14:58:47 ON 16 MAR 2006
1.1
                STRUCTURE UPLOADED
                S L1
     FILE 'REGISTRY' ENTERED AT 14:59:34 ON 16 MAR 2006
L2
           3981 S L1 FULL
     FILE 'CAPLUS' ENTERED AT 14:59:35 ON 16 MAR 2006
          13736 S L2 FULL
L3
            701 S L3 AND ( QUATERNARY AMMONIUM OR QUATERNARY PHOSPHONIUM)
L4
L5
             41 S L4 AND DOCUSATE
             23 S L5 AND PY<2002
L6
              2 S L6 AND SULFOSUCCINIC ACID
L7
                STRUCTURE UPLOADED
L8
                S L8
     FILE 'REGISTRY' ENTERED AT 15:21:49 ON 16 MAR 2006
L9
            204 S L8 FULL
     FILE 'CAPLUS' ENTERED AT 15:21:50 ON 16 MAR 2006
L10
            295 S L9 FULL
L11
            224 S L10 AND PY<2002
L12
             10 S L11 AND COMPOSITION
L13
              5 S L11 AND (QUATERNARY AMMONIUM OR QUATERNARY PHOSHONIUM)
=> s lll and liquid
        714766 LIQUID
L14
            11 L11 AND LIQUID
=> d 1-11, ibib abs hitstr
L14 ANSWER 1 OF 11 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
                         2001:409397 CAPLUS
DOCUMENT NUMBER:
                         135:220343
TITLE:
                         Analysis of sulfophthalimide and some of its
                         derivatives by liquid chromatography-
                         electrospray ionization tandem mass spectrometry
                         Reemtsma, T.
AUTHOR (S):
CORPORATE SOURCE:
                         Department of Water Quality Control, Technical
                         University of Berlin, Berlin, D-10623, Germany
SOURCE:
                         Journal of Chromatography, A (2001), 919(2),
                         289-297
                         CODEN: JCRAEY; ISSN: 0021-9673
PUBLISHER:
                         Elsevier Science B.V.
DOCUMENT TYPE:
                         Journal
LANGUAGE:
                         English
     A system was developed for the separation of sulfophthalimide (SPI),
     sulfophthalamide (SPAM), sulfophthalamic acid (SPAA) and sulfophthalic
     acid (SPA) by ion-pair liquid chromatog. and their detection by electrospray
     ionization tandem mass spectrometry (ESI-MS-MS). Except for SPAM, the 3-
     and 4-sulfo-isomers of the analytes were separated by HPLC using volatile
     tributylamine as ion-pairing agent. Initial fragmentations of the
     analytes in the neg. mode involve losses of CO2 or HNCO or condensation
     via H2O or NH3 elimination. Ortho-effects of the sulfonate group were
     recognized in the fragmentation of the resp. 3-sulfo-isomers and allowed
     the authors to assign the order of elution of the SPAA isomers. Quant.
     anal. of these sulfonated aromatic compds. with MRM detection was elaborated
     and resulted in detection limits ranging from 1 pg for SPA to 13 pg for
     SPAA isomers and in limits of quantification of 2-10 \mu g/L for 5 \mu L
     vols. of injected tap water, municipal wastewater or industrial effluents
     up to salt contents of 0.5-1 g/L. The method was applied to study the
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isomer-specific chemical and microbial transformations of SPI, which was previously shown to be formed by white-rot fungi from sulfophthalocyanine

textile dyes.

IT 89-08-7, 4-Sulfophthalic acid 67892-43-7, 3-Sulfophthalic acid

RL: ANT (Analyte); ANST (Analytical study)

(analyte; anal. of sulfophthalimide and some of its derivs. by liquid chromatog.-electrospray ionization tandem mass spectrometry)

RN 89-08-7 CAPLUS

1,2-Benzenedicarboxylic acid, 4-sulfo- (9CI) (CA INDEX NAME)

CN

CN

RN 67892-43-7 CAPLUS

1,2-Benzenedicarboxylic acid, 3-sulfo- (9CI) (CA INDEX NAME)

REFERENCE COUNT: 25 THERE ARE 25 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L14 ANSWER 2 OF 11 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1999:699113 CAPLUS

DOCUMENT NUMBER:

131:300576

TITLE:

Sulfonated porphyrazine dyes, and ink-jet inks, color

filters, liquid crystal panels, and

computers using them

INVENTOR(S):

Hirose, Masashi; Kashiwazaki, Akio; Shirota,

Kachihiro; Nakazawa, Koichiro; Yamashita, Yoshihisa;

Yokoyama, Mayumi

PATENT ASSIGNEE(S):

Canon K. K., Japan

SOURCE:

Jpn. Kokai Tokkyo Koho, 22 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 11302285 PRIORITY APPLN. INFO.: OTHER SOURCE(S): GI	A2	19991102	JP 1998-111170	19980421 <
	MARPAT	131:300576	JP 1998-111170	19980421

The dyes comprise I [A1-A4 = (substituted) (N-containing hetero) aromatic ring; M = 2H, divalent metal, tri- or tetravalent metal derivative; D = alkali metal, NH4; m = 1-4; n = 0-3; m + n = 1-4]. Thus, reaction of pyridine-2,3-dicarboxylic acid with urea and CuCl2 in the presence of ammonium molybdate gave porphyrazine, which was sulfonated and neutralized by NaOH. A glass substrate having black matrixes was coated with N-methylolacrylamide-hydroxyethyl methacrylate (1:1) copolymer, printed using an ink containing the sulfonated porphyrazine and C.I. Direct Blue 199, and covered with SS 7625 (acrylic thermosetting resin) to give a color filter showing good transparency, heat and light resistance, and printing precision.

89-08-7, 4-Sulfophthalic acid

RL: RCT (Reactant); RACT (Reactant or reagent)

(in preparation of sulfonated porphyrazine dyes for ink-jet inks for color filters for liquid crystal panels for computers)

89-08-7 CAPLUS

CN 1,2-Benzenedicarboxylic acid, 4-sulfo- (9CI) (CA INDEX NAME)

IT

RN

L14 ANSWER 3 OF 11 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1996:200943 CAPLUS

DOCUMENT NUMBER: 124:305734

TITLE: High-sensitivity conductivity detection in

nonsuppressed ion chromatography using

sulfoisophthalic acid as eluent

AUTHOR(S): Watanabe, Hideki; Yokoyama, Yukio; Sato, Hisakuni

CORPORATE SOURCE: Laboratory of Analytical Chemistry, Faculty of

Engineering, Yokohama National University, Tokiwadai

156, Hodogaya-ku, Yokohama, 240, Japan

SOURCE: Journal of Chromatography, A (1996), 727(2),

311-16

CODEN: JCRAEY; ISSN: 0021-9673

PUBLISHER: Elsevier
DOCUMENT TYPE: Journal
LANGUAGE: English

AB A nonsuppressed ion chromatog. (IC) system for the high-sensitivity detection of common anions was developed using sulfoisophthalic acid as the eluent. The detection sensitivity was ten times higher than that using conventional nonsuppressed IC with sodium phthalate as eluent, and was almost the same as that using conventional suppressed IC with a carbonate-hydrogen carbonate eluent under the same elec. conditions with a

conductivity detector. Temperature regulation was very important in minimizing the baseline drift. A com. incubator, in which a separation column and a sample injector were placed, was useful. The developed nonsuppressed system facilitated the determination of low concns. of phosphate, chloride, bromide, nitrate and sulfate at micromolar levels.

89-08-7, 4-Sulfophthalic acid IT

RL: ARU (Analytical role, unclassified); NUU (Other use, unclassified); ANST (Analytical study); USES (Uses)

(as eluent in nonsuppressed ion chromatog. of anions in comparison to sulfoisophthalic acid as eluent)

89-08-7 CAPLUS RN

1,2-Benzenedicarboxylic acid, 4-sulfo- (9CI) (CA INDEX NAME)

SO<sub>3</sub>H со2н CO2H

CN

L14 ANSWER 4 OF 11 CAPLUS COPYRIGHT 2006 ACS on STN

KIND

ACCESSION NUMBER: 1992:96489 CAPLUS

DOCUMENT NUMBER: 116:96489

TITLE: Graft copolymers from poly(arylene sulfide) backbones

and liquid crystalline side chains

INVENTOR(S): Koehler, Burkhard; Wehrmann, Rolf; Pielartzik, Harald;

Heinz, Hans Detlef; Ebert, Wolfgang

APPLICATION NO.

DATE

PATENT ASSIGNEE(S): Bayer A.-G., Germany

SOURCE: Ger. Offen., 7 pp.

CODEN: GWXXBX

DATE

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.

	DE 3940793	A1	19910613	DE 1989-3940793	19891209 <
	EP 432561	A2	19910619	EP 1990-122583	19901127 <
	EP 432561	A3	19911121		
	EP 432561	B1	19950607		
	R: BE, DE, FR,				
	JP 03250024	A2	19911107	JP 1990-407411	19901208 <
PRIO				DE 1989-3940793 A	
AB	The title copolymer	s are f	ormed from c	arboxyl group-, dicarbox	xylic acid
	anhydride group-, h	ydroxy (	group-, and/	or amino group-modified	poly(arylene
	sulfide) backbones,	produc	ed by reacti	ng sulfonic acid group-	or nitro
	group containing ar	omatic	compds. with	a poly(arylene sulfide)	at temps, above
	the m.p. of the pol	y(aryle:	ne sulfide),	which are reacted with	liquid crystalline
	polyester side chai	n mater	ials under c	onditions which result :	in the
	formation of covale	nt bond	s between th	e side chains and the ba	ackbone.
IT	89-08-7DP, 4-Sulfop	hthalic	acid, react	ion products with	
	polyparaphenylene s	ulfides	, graft poly	mers with liquid crystal	lline polvesters
	RL: SPN (Synthetic	prepara	tion): PREP	(Preparation)	rime perfectes
	(preparation of)			,	
RN	89-08-7 CAPLUS				

1,2-Benzenedicarboxylic acid, 4-sulfo- (9CI) (CA INDEX NAME) CN

INVENTOR(S):

L14 ANSWER 5 OF 11 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1991:516858 CAPLUS

DOCUMENT NUMBER: 115:116858

TITLE: Stable thickened liquid cleaning composition

containing bleach
Wise, Rodney Mahlon

PATENT ASSIGNEE(S): Procter and Gamble Co., USA

SOURCE: Eur. Pat. Appl., 16 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 421738	A2	19910410	EP 1990-310787	19901002 <
EP 421738	A3	19911016		
EP 421738	B1	19960522		
R: AT, BE, CH,	DE, DK	, ES, FR, GE	B, GR, IT, LI, LU, NL,	SE
CA 2026332	AA	19910405	CA 1990-2026332	19900927 <
CA 2026332	C ·	19950221		
AT 138410	E	19960615	AT 1990-310787	19901002 <
ES 2087132	T3	19960716	ES 1990-310787	19901002 <
AU 9063786	A1	19910411	AU 1990-63786	19901003 <
AU 648993 ·	B2	19940512		
JP 03166299	A2	19910718	JP 1990-266172	19901003 <
JP 2766064	B2	19980618		
US 5169552	A	19921208	US 1991-708826	19910529 <
PRIORITY APPLN. INFO.:			US 1989-417123	A 19891004
OMITED COLLDGE (C)	****			<del></del>

OTHER SOURCE(S): MARPAT 115:116858

The title composition, useful for automatic dishwashing and hard surface cleaning, contains Cl bleach, crosslinked polymer containing carboxy groups, buffering agent to give pH >10 and rheol. stabilizing agent selected from BzOH, BzOH substituted by 1-3 CO2H, Cl, Br, SO3H, NO2, OMe, or Cl-4 alkyl groups, and their alkali metal salts. An automatic dishwashing composition contained Na5P3O10 4.67, Na4P2O7 12.60, Na silicate 3.27, K2CO3 3.91, Na2CO3 2.61, available Cl (as NaOCl) 0.93, KOH 0.84, monostearyl acid phosphate 0.03, acrylic acid polymer (Sokalan PHC 25) 1.07, Al2O3 (as Na aluminate) 0.03, and BzOH 0.47%, the balance being water, perfume, dye, and KOH (to give pH 12.2-12.3).

IT 89-08-7, 4-Sulfophthalic acid

RL: USES (Uses)

(rheol. stabilizer, in liquid cleaner containing chlorine bleach)

RN 89-08-7 CAPLUS

1,2-Benzenedicarboxylic acid, 4-sulfo- (9CI) (CA INDEX NAME)

CN

L14 ANSWER 6 OF 11 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1991:484452 CAPLUS

DOCUMENT NUMBER: 115:84452

TITLE: Ion-exchange chromatographic determination of anions

by indirect photometric detection: comparison of eluent ions with respect to sensitivity enhancement Motomizu, Shoji; Oshima, Mitsuko; Hironaka, Takashi

CORPORATE SOURCE: Fac. Sci., Okayama Univ., Okayama, 700, Japan SOURCE: Analyst (Cambridge, United Kingdom) (1991),

116(7), 695-700

CODEN: ANALAO; ISSN: 0003-2654

DOCUMENT TYPE: Journal LANGUAGE: English

Aromatic sulfonate and carboxylate eluent ions were examined for use in the sensitive determination of inorg. anions by indirect photometric ion chromatog. The naphthalene-1,3,6-trisulfonate ion was found to be the most sensitive for use as the eluent ion, the detection limit being as low as 1 + 10/-8 mol dm-3. The naphthalene-1,5-disulfonate ion is recommended for the anal. of water samples containing anions at concns. of between 1 + 10-6 and 1 + 10-5 mol dm-3. These two eluent ions have several advantages over other choices: (i) detection is carried out at longer wavelengths (near 300 nm); (ii) the eluent ions are easily soluble in water and subsequently stable; (iii) their elution strength is not influenced by pH change; (iv) the eluent ions do not form any metal complexes; and (v) the reagents are inexpensive and com. available.

IT 46687-30-3

AUTHOR (S):

RL: ANST (Analytical study)

(as eluent for ion-exchange chromatog. determination of anions with indirect photometric detection)

46687-30-3 CAPLUS

CN 1,2-Benzenedicarboxylic acid, 4-sulfo-, ion(3-) (9CI) (CA INDEX NAME)

RN

L14 ANSWER 7 OF 11 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1989:596981 CAPLUS

DOCUMENT NUMBER: 111:196981

TITLE: Oil-based liquid recording compositions for

ink-jet printing

INVENTOR(S): Tanaka, Mitsugi; Sakai, Takeo
PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan
SOURCE: Jpn. Kokai Tokkyo Koho, 7 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 01020278	A2	19890124	JP 1987-174497	19870713 <
JP 05079268	B4	19931101		
CORITY APPLN. INFO.:			JP 1987-174497	19870713

OTHER SOURCE (S): MARPAT 111:196981

The storage-stable title compns., giving light- and water-resistant images, contain MPc(SO2NHR)n (I; M = metal; Pc = phthalocyanine residue; R = alkyl which has a tertiary C linked to an N atom; n = 1-4). I (M = Cu, R = CMe2CH2CHMe2, n = 4) 5, di-Et phthalate 30, diisopropyl adipate 45,

and N,N-diethyldodecanamide 20 parts were mixed and filtered to give a title composition, which was jet-printed onto a silica- and poly(vinyl alc.)-coated paper, giving clear images showing ≤1% degradation of color d. after a 3-mo indoor exposure.

33562-89-9 IT

RL: USES (Uses)

(dyes from, for oil-based inks., for ink-jet printing, with improved durability)

RN33562-89-9 CAPLUS

1,2-Benzenedicarboxylic acid, 4-sulfo-, monosodium salt (9CI) (CA INDEX CN NAME)

# **N**a

AUTHOR (S):

CORPORATE SOURCE:

L14 ANSWER 8 OF 11 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1989:95183 CAPLUS

DOCUMENT NUMBER: 110:95183

TITLE: Biological activities of phthalocyanines. X.

Syntheses and analyses of sulfonated phthalocyanines Ali, Hasrat; Langlois, Rejean; Wagner, J. Richard; Brasseur, Nicole; Paquette, Benoit; Van Lier, Johan E.

Fac. Med., Univ. Sherbrooke, Sherbrooke, QC, J1H 5N4,

Can.

SOURCE: Photochemistry and Photobiology (1988),

47(5), 713-17

CODEN: PHCBAP; ISSN: 0031-8655

DOCUMENT TYPE: Journal LANGUAGE: English

Synthetic methods to obtain selectively sulfonated metallophthalocyanines are compared. Both condensation [of 3,4-(HO2C)2C6H3SO3Na with o-(HO2C)2C6H4 derivs., H2NCONH2, and metal salts] and direct sulfonation (of metallophthalocyanines with SO2-H2SO4) procedures lead to mixts. of mono- to tetrasulfonated products which are resolved by reversed-phase liquid chromatog. in buffered H2O-MeOH. The proportion of sulfonated derivs. is examined as a function of the starting reagents for the condensation method, and as a function of the temperature and reaction time for the direct sulfonation procedure. The number of SO3H groups per phthalocynine mol. is determined by oxidative degradation of the sulfonated phthalocyanine ring followed by quant. chromatog. anal. of the sulfophthalimide and phthalimide fragments. IT

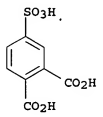
33562-89-9

RN

RL: RCT (Reactant); RACT (Reactant or reagent) (cyclocondensation reactions of, with phthalic acid derivs., urea, and metal salts, phthalocyanine derivs. from)

33562-89-9 CAPLUS

1,2-Benzenedicarboxylic acid, 4-sulfo-, monosodium salt (9CI) (CA INDEX CN NAME)



Na

L14 ANSWER 9 OF 11 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1985:39213 CAPLUS

DOCUMENT NUMBER: 102:39213

TITLE: Separation of sulfonate and carboxylate mixtures by

ion-exchange high-performance liquid

chromatography

AUTHOR(S): Bear, G. R.; Lawley, C. W.; Riddle, R. M.

CORPORATE SOURCE: Expl. Prod. Serv. Dep., Texaco Inc., Bellaire, TX,

77401, USA

SOURCE: Journal of Chromatography (1984), 302, 65-78

CODEN: JOCRAM; ISSN: 0021-9673

DOCUMENT TYPE: Journal LANGUAGE: English

Aromatic sulfonate and carboxylate mixts. were separated by an ion-exchange high-performance liquid chromatog. method. The separation was carried out on a strong anion exchanger (quaternary amine) with a gradient of 3 solvents: THF-water (50:50), THF-0.1M KH2PO4 (pH 4.5) (50:50), and THF-0.2M KH2PO4 (pH 6.5) (50:50). The change in ionic strength and pH of the mobile phase during elution resulted in excellent resolution of mixts. by charge and ionic group. Small variations in retention time within each class of ionic group were noted and are due to electronic and steric effects introduced by substituents on the hydrophobic part of the mol. When applied to petroleum sulfonates, i.e., complex mixts. of alkylaryl sulfonates, this procedure gives information on the degree of sulfonation as well as the extent of variation in the structure of the alkylaryl part of the anions.

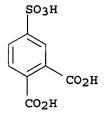
IT 89-08-7

RL: ANST (Analytical study)

(anion-exchange HPLC of, retention in)

RN 89-08-7 CAPLUS

CN 1,2-Benzenedicarboxylic acid, 4-sulfo- (9CI) (CA INDEX NAME)



L14 ANSWER 10 OF 11 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1978:492182 CAPLUS

DOCUMENT NUMBER: 89:92182

TITLE: Mercaptan oxidation in a liquid hydrocarbon

with a metal phthalocyanine catalyst

INVENTOR(S): Douglas, Walter M. PATENT ASSIGNEE(S): UOP Inc., USA

PATENT ASSIGNEE(S): UOP Inc., USA SOURCE: U.S., 7 pp. CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: PATENT · INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 4088569	Α	19780509	US 1977-817872	19770721 <
US 4049572	Α	19770920	US 1976-660899	19760224 <
PRIORITY APPLN. INFO.:			US 1976-660899 A3	3 19760224
			US 1977-787756 A3	3 19770421

AB A Co phthalocyaninesulfonate [30638-08-5] catalyst for kerosine sweetening was prepared by reaction of 4-sulfophthalic acid 89-08-7], CoSO4, ammonium molybdate, urea, and water, addition of the mixture to phthalic anhydride [85-44-9] and heating at 190-215° for 3 h and to 260-70° for 3.5 h.

IT 89-08-7

> RL: RCT (Reactant); RACT (Reactant or reagent) (reaction of, with cobalt sulfate, phthalic anhydride, and urea, in manufacture of oxidation catalysts)

89-08-7 CAPLUS RN

1,2-Benzenedicarboxylic acid, 4-sulfo- (9CI) (CA INDEX NAME)

L14 ANSWER 11 OF 11 CAPLUS COPYRIGHT 2006 ACS on STN

1933:59141 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 27:59141 ORIGINAL REFERENCE NO.: 27:5315d-h

TITLE: Mercury as a sulfonation catalyst

AUTHOR (S): Lauer, Karl

SOURCE: Journal fuer Praktische Chemie (Leipzig) (1933

), 138, 81-91

CODEN: JPCEAO; ISSN: 0021-8383

DOCUMENT TYPE: Journal LANGUAGE: Unavailable

The p-directing action of Hg is shown in the following expts. PhNO2, with 20% oleum at 90°, gives 3% p- and 97% m-SO3H derivative; in the presence of 5% Hg, there results 25% p- and 75% of the m-isomer. with 10% oleum at 150°, gives 14% p- and 86% m-SO3H derivative; with 5% Hg, there results 5% o-, 26% p- and 69% m-isomers. PhSO3H with 20% oleum at 200° gives 5% p- and 95% m-SO3H derivs.; 5% Hg gives 31% p- and 69% m-isomers. The behavior of HgCl compds. with H2SO4 was also studied, using 92% H2SO4 and 3% SO3; o-HOC6H4HgCl gave 38% o- and 62% p-SO3H derivative and 93 and 7%, resp.; p-HOC6H4HgCl gave 41 and 59%, 6 and 94% o- and p-SO3H derivs., resp. p-MeC6H4HgCl gives 29 and 71%, and 5 and 95% o- and p-SO3H derivs., resp. o-O2NC6H4HgCl and 92% H2SO4 give 5% p- and 95% m-SO3H derivs.; with 20% SO3 there results 94% of the o- and 6% of the m-SO3H derivs. o-HO2CC6H4HgCl and 92% H2SO4 give 8% p- and 92% m-SO3H derivs.; 10% SO3 gives 97% o- and 3% m-SO3H derivs. o-C6H4Me2 with 0, 2, and 10% Hg gives, resp., 0 and 100, 8 and 92, and 22 and 78% of the 3- and 4-SO3H derivs. o-C6H4Cl2, with 0, 2 and 10% Hg, gives, resp., 0 and 100, 16 and 84, 26 and 74% of the 3- and 4-SO3H derivs. o-C6H4Br2 with 0 and 10% Hg, gives 0 and 100, and 24 and 76% of the 3- and 4-SO3H derivs. o-C6H4(CO2H)2, with 0 and 5% Hg, gives 0 and 100, 50 and 50% of the 3- and 4-SO3H derivs. 3,5-Disulfophthalic acid (I) is formed in 50% yield from C6H4(CO)2O with Hg and oleum; 46% of the 4-SO3H derivs., is also formed. 3-Sulfophthalic acid gives 85-9% of I. The 4-isomer is not further sulfonated. Na o-xylene-3-sulfonate seps. with 1 mol. H2O, the di-Cl derivative with 2 mols. H2O and the di-Br derivative with 1 mol. H2O.

(preparation of)
216451-89-7 CAPLUS
1,2-Benzenedicarboxylic acid, 3,5-disulfo- (9CI) (CA INDEX NAME)

RN

CN